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PE CONTENT TERRET

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Cnfrm. No. : 7683

Filed: January 19, 2001

For : HYPERSENSITIVE RESPONSE INDUCED

RESISTANCE IN PLANTS BY SEED

TREATMENT

DECLARATION OF ZHONG-MIN WEI UNDER 37 C.F.R. § 1.132

I, ZHONG-MIN WEI, pursuant of 37 C.F.R. § 1.132, declare:

- 1. I received a B.S. degree in Biology from Zhejiang University, Zhejiang, China in 1982, an M.S. degree in Plant Pathology from Nanjing Agricultural University, Nanjing, China in 1984, and a Ph.D. degree in Molecular Biology from Nanjing Agricultural University and Academy of Science, Shanghai, China in 1987.
- 2. I am currently employed as Chief Scientific Officer and Vice President of Research and Development at EDEN Bioscience Corporation in Bothell, Washington.
 - 3. I am an inventor of the above-identified application.
- 4. I am presenting this declaration to show that hypersensitive response elicitors from a diverse range of plant pathogenic bacteria (1) are an art-recognized class of proteins where results achieved with one such protein would be expected when other proteins in this class are used and (2) share the unique ability to cause distinct plant responses. Specifically, treatment of a variety of plants and plant seeds with hypersensitive response elicitors was shown to induce plant disease resistance as compared with plants and plant seeds not treated with a hypersensitive response elicitor; and transgenic expression of hypersensitive response elicitors in transgenic plants was shown to induce plant disease resistance as compared to null transfected plants or wild-type plants.

- 5. In plants, the hypersensitive response phenomenon results from an incompatible interaction between plant pathogens and non-host plants. As explained in Gopalan et al., "Bacterial Genes Involved in the Elicitation of Hypersensitive Response and Pathogenesis," Plant Disease 80: 604-10 (1996) ("Gopalan") (attached hereto as Exhibit 1), these types of interactions involve, for example, a bacterial plant pathogen attempting to infect a host plant, and the host plant preventing proliferation of the pathogen by the collapse and death, or necrosis, of plant leaf cells at the site of infection. This is distinct from a compatible interaction between a bacterial plant pathogen and a host plant in which the bacteria is capable of proliferation, resulting in the spread of the pathogen throughout the plant and the manifestation of disease symptoms. Id. at 604.
- 6. Hypersensitive response elicitors within a given genus are often homologous to elicitors from different pathogenic species and strains of the same genus. For example, homologs of hypersensitive response elicitors from *Erwinia amylovora* and *Pseudomonas syringae* have been identified in different bacteria species and strains from the genera *Erwinia* and *Pseudomonas*, respectively. *See* Gopalan.
- 7. In addition, numerous reported studies confirm that a gene encoding a hypersensitive response elicitor from a particular source genus can be used to isolate a corresponding hypersensitive response elicitor gene from different species and strains of that same genus. For example, in Bauer et al., "Erwinia chrysanthemi Harpin_{Ech}: An Elicitor of the Hypersensitive Response that Contributes to Soft-Rot Pathogenesis," MPMI 8(4): 484-91 (1995) ("Bauer") (attached hereto as **Exhibit 2**), the Erwinia amylovora hypersensitive response elicitor encoding gene was used as a probe to isolate, clone, and sequence the gene encoding the Erwinia chrysanthemi hypersensitive response elicitor, as follows:

The cosmids were probed in colony blots with a 1.3-kb *Hind*III fragment from pCPP1084, which contains the *E. amylovora hrpN* gene (Wei et al. [, "Harpin Elicitor of the Hypersensitive Response Produced by the Plant Pathogen *Erwinia amylovora*," Science 257:85-88 (]1992[)]). pCPP2157, one of the three cosmids hybridizing with the probe, was digested with several restriction enzymes, and the location of the *hrpN_{Ech}* gene in those fragments was determined by probing a Southern blot with *E. amylovora Hind*III fragment. Two fragments, each containing the entire *hrpN_{Ech}* gene, were subcloned into different vectors: pCPP2142 contained an 8.3-kb *Sal*I fragment in pUC119 (Vieira and Messing [,"Production of Single-Stranded Plasmid DNA," Methods Enzymol., 153:3-11(] 1987[)]), and pCPP2141 contained a 3.1-kb *Pst*I fragment in pBluescript II SK(-) (Stratagene, La Jolla, CA).

Sequence of hrpN_{Ech}

The nucleotide sequence of a 2.4-kb region of pCPP2141 encompassing $hrpN_{Ech}$ was determined. The portion of that sequence extending from the putative ribosome-binding site through the $hrpN_{Ech}$ coding sequence to a putative rhoindependent terminator is presented in Figure 1.

See page 485.

8. In the same manner as described in Bauer *supra*, Cui et al., "The RsmA Mutants of *Erwinia carotovora* subsp. *carotovora* Strain Ecc71 Overexpress *hrp*N_{Ecc} and Elicit a Hypersensitive Reaction-like Response in Tobacco Leaves," *MPMI* 9(7): 565-73 (1996) ("Cui") (attached hereto as **Exhibit 3**) demonstrates that the gene encoding the *Erwinia carotovora* hypersensitive response elicitor can be isolated, sequenced, and cloned by using the *Erwinia chrysanthemi* hypersensitive response elicitor encoding gene to probe the genomic library of *Erwinia carotovora*. Further, Cui (at page 572) states the following:

The genomic library of *E. carotovora* subsp. *carotovora* strain Ecc71 in pLARF5 was screened by in situ colony hybridization with a 0.75-kb internal *ClaI* fragment of *hrpN* of *E. chrysanthemi* (Bauer et al.[, "*Erwinia chrysanthemi* Harpin_{Ech}: An Elicitor of the Hypersensitive Response that Contributes to Soft-Rot Pathogenesis," MPMI 8(4): 484-91 (]1995). Two cosmids, pAKC921 and pAKC922, that hybridized with the probe were isolated. The subclones (pAKC923 and pAKC924, Table 1) carrying *hrpN* DNA were used for sequence analysis.

- 9. The gene encoding the hypersensitive response elicitor of *Erwinia* amylovora has also been used as a probe to isolate and clone the gene encoding the hypersensitive response elicitor of *Erwinia stewartii*. It was found that antibodies raised against the hypersensitive response elicitor of *Erwinia stewartii* cross-reacted with the hypersensitive response elicitor of *Erwinia amylovora*. See Ahmad et al., "Harpin Is Not Necessary for the Pathogenicity of Maize," 8th Int'l Cong. Molec. Plant Microbe Inter. July 14-19, 1996 ("Ahmad") (attached hereto as **Exhibit 4**).
- 10. The genes encoding the HrpN hypersensitive response elicitor from several strains of *Erwinia pyrifolia* have since been cloned. As reported in Jock et al., "Molecular Differentiation of *Erwinia amylovora* Strains from North America and of Two Asian Pear Pathogens by Analyses of PFGE Patterns and *hrpN* genes," *Environ. Microbiol.* 6(5): 480-490 (2004) ("Jock") (attached hereto as **Exhibit 5**), the *hrpN* genes were amplified with PCR consensus primers that were deduced by comparison of the known nucleotide

sequences of Erwinia amylovora hrpN and Erwinia stewartii hrpN. Indeed, Jock (at page 489) recites the following:

Erwinia pyrifoliae and the Erwinia strains from Japan were considered to be sufficiently related to E. amylovora to amplify their genes with the Erwinia PCR consensus primers given above. This was indeed possible and allowed cloning and sequencing of their hrpN DNA fragments. . . .

- from the genus *Pseudomonas*. An internal fragment of the hypersensitive response elicitor from *Pseudomonas syringae* pv. *syringae* (i.e., *hrpZ*) was used to identify and isolate the hypersensitive response elicitors from *P. syringae* pv. *glycinea* and *P. syringae* pv. *tomato*. Significant amino acid sequence similarities were identified between the various *Pseudomonas syringae* elicitors. *See* Preston et al., "The HrpZ Proteins of *Pseudomonas syringae* pvs. *syringae*, *glycinea*, and *tomato* Are Encoded by an Operon Containing *Yersinia ysc* Homologs and Elicit the Hypersensitive Response in Tomato But Not Soybean," *MPMI* 8(5): 717-32 (1995) ("Preston") (attached hereto as **Exhibit 6**).
- within the *hrp* gene cluster or proximate to the *hrp* gene cluster in *hrp* regulons. For example, *hrpN* from *Erwinia amylovora* was located within the *hrp* gene cluster, as was *hrpZ* from *Pseudomonas syringae*. The *popA* gene, encoding a hypersensitive response elicitor from *Pseudomonas solanacearum*, was located on the left flank of the *hrp* gene cluster within a *hrp* regulon. *See* Bonas, "*hrp* Genes of Phytopathogneic Bacteria," *Current Topics in Microbiology and Immunology* 192: 79-98 (1994) ("Bonas I") (attached hereto as **Exhibit 7**) and Alfano et al., "The Type III (Hrp) Secretion Pathway of Plant Pathogenic Bacteria: Trafficking Harpins, Avr Proteins, and Death," *Journal of Bacteriology* 179: 5655-5662 (1997) ("Alfano") (attached hereto as **Exhibit 8**). Similar to the *popA* gene, *hreX*, the gene encoding the hypersensitive response elicitor from *Xanthomonas campestris*, was located on the left flank of the *hrp* gene cluster. *See* Swanson et al., "Isolation of the *hreX* Gene Encoding the HR Elicitor Harpin (Xcp) from *Xanthomonas campestris* pv. *pelargonii*," *Phytopathology* 90: s75 (1999) ("Swanson") (attached hereto as **Exhibit 9**).
- 13. The characteristics that distinguish hypersensitive response elicitors as a distinct class of molecules are clearly apparent when considering the different elicitors' secretion mechanisms, regulation, biochemical characteristics, and biological activities.

- shown to be secreted through the type III, *hrp* dependent secretion pathway. The type III secretion pathway is a highly conserved and unique mechanism for the delivery of pathogenicity related molecules in gram-negative bacteria. The *hrp* gene cluster is largely composed of components of the type III secretion system. *See* Bogdanove et al., "Unified Nomenclature for Broadly Conserved *hrp* Genes of Phytopathogenic Bacteria," *Molec. Microbiol.* 20:681-83 (1996) (attached hereto as **Exhibit 10**); and Alfano.
- subsequently the genes encoding the components of the type III secretion system and hypersensitive response elicitors, is controlled by environmental factors. Specifically, transcriptional expression of these genes is induced under conditions that mimic the plant apoplast, such as low concentrations of carbon and nitrogen, low temperature, and low pH. See Wei et al., "Regulation of hrp Genes and Type III Protein Secretion in *Erwinia* amylovora by HrpX/HrpY, a Novel Two-Component System, and HrpS," MPMI 13(11): 1251-1262 (2000) ("Wei I") (attached hereto as **Exhibit 11**); and Bonas I.
- 16. Biochemically, hypersensitive response elicitors have a number of common characteristics. These include being glycine rich, heat stable, hydrophilic, lacking of an N-terminal signal sequence, and susceptible to proteolysis. *See* Bonas, "Bacterial Home Goal by Harpins," *Trends Microbiol* 2: 1-2 (1994) (attached hereto as **Exhibit 12**); Bonas I; Gopalan; and Alfano.
- 17. In addition, hypersensitive response elicitors share a unique secondary structure that has been associated with these elicitors' distinct biological activities (described below). The structure has two primary components, an alpha helix unit and a relaxed acidic unit having a sheet or random turn structure. In the absence of one or both of these components, hypersensitive response elicitation does not occur. *See* WO 01/98501 to Fan et al. (attached hereto as **Exhibit 13**).
- 18. In addition to eliciting the hypersensitive response in a broad range of plant species, as explained by Wei et al., "Harpin from *Erwinia amylovora* Induced Plant Resistance," *Acta Horticulture* 411: 223-225 (1996) ("Wei II") (attached hereto as **Exhibit** 14) and by Alfano, hypersensitive response elicitors also share the ability to induce specific plant responses. The induction of plant disease resistance, plant growth enhancement, and plant stress resistance are three plant responses that result from treatment of plants or plant seeds with a hypersensitive response elicitor from a gram-negative plant pathogen.

- 19. As described in Wei II, treatment of plants with the hypersensitive response elicitor HrpN from *Erwinia amylovora* resulted in disease resistance to a broad range of plant pathogens. For example, HrpN induced disease resistance to southern bacterial wilt (*Pseudomonas solanacearum*) in tomato, tobacco mosaic virus in tobacco, and bacterial leaf spot (*Gliocladium cucurbitae*) in cucumber.
- 20. The hypersensitive response elicitor HrpZ from *Pseudomonas* syringae was reported to induce disease resistance in cucumber to a diverse range of pathogens, including the fungal disease *Colletotrichum lagenarium*, tobacco necrosis virus, and bacterial angular leaf spot (*Pseudomonas syringae* pv. *lachrymans*). See Strobel et al., "Induction of Systemic Acquired Resistance in Cucumber by *Pseudomonas syringae* pv. syringae 61 HrpZ_{Pss} Protein," *Plant Journal* 9(4): 431-439 (1996) (attached hereto as **Exhibit** 15).
- 21. The hypersensitive response elicitor HrpZ from *Pseudomonas* syringae was reported to induce disease resistance in transgenic tobacco to powdery mildew (*Erysiphe cichoracearum*), and in transgenic rice to rice blast fungus (*Magneporthe grisea*). See U.S. Patent Application Publ. No. 2004/0073970 to Takakura et al. (attached hereto as **Exhibit 16**) at Example 4. The HrpZ-expressing transgenes included transgenes with either a weak or a strong constitution promoter, an inducible promoter, or a tissue-specific promoter. *Id.* at Example 3.

Hypersensitive Response Elicitors Induce Plant Disease Resistance

- 22. As demonstrated by the following experimental evidence in paragraphs 23 and 24 below, treatment of tomato and tobacco plants with the hypersensitive response elicitor HreX from *Xanthomonas campestris* pv. *pelargonii* induced disease resistance in the plants against bacterial wilt and tobacco mosaic virus.
- Caused by the pathogenic bacterium *Pseudomonas solanacearum* K_{60}) was investigated as follows. Approximately 30 days after sowing, tomato plants were sprayed with either a dilution of HreX or 5 mM potassium phosphate buffer, pH 6.8 (the same buffer used to dilute the HreX solution). Six days after treatment, inoculation was performed by slicing the soil of the pot containing the tomato plant 4 times and applying 40 ml of solution containing 1 x 10^6 colony forming units ("cfu") per ml of *P. solanacearum* K_{60} to the soil. Disease severity ratings were recorded at 7, 9, and 13 days after inoculation ("DAI"), as shown below in

Table 1. As these results demonstrate, plants treated with HreX exhibited substantially more disease resistance than the buffer-treated control plants.

Table 1. Pseudomonas solanacearum Disease Resistance from Treatment of Tomato with HreX.

Treatment Groups ^a	Disease Index (7 DAI)	Disease Index (9 DAI)	Disease Index (12 DAI)	% Difference (12 DAI)
HreX	0.12	0.22	0.22	38.89
Buffer	0.16	0.3	0.36	na

Each group consisted of 1 pot containing 10 plants.

24. Experiments examining the induction of systemic disease resistance in tobacco from treatment with HreX were conducted as follows: Diluted HreX was sprayed on all but the bottom most full-sized leaf of six- to eight-week-old tobacco plants (Xanthi). The bottom most full-sized leaf was covered during spraying so as not to receive residual spray. Three days after the spray treatment, the unsprayed leaf and the leaf opposite it, were lightly dusted with diatomaceous earth. Thereafter, 20 μl of a 1.7 μg/ml solution of tobacco mosaic virus ("TMV") was applied to both leaves dusted with diatomaceous earth. The TMV was gently and evenly spread across the leaves. Approximately 5 minutes after inoculation, the plants were lightly rinsed to remove the diatomaceous earth. Three days after inoculation, the number of TMV lesions on the unsprayed and sprayed leaves for each plant was recorded, as shown below in Table 2. As these results demonstrate, plants treated with HreX exhibited substantially more disease resistance than the buffer-treated control plants.

Table 2. Tobacco Mosaic Virus Resistance in Tobacco from Treatment with HreX.

				Nu	mber of TMV	Lesions	on Leaf			
Treatment	Treated leaves			Untreated leaves						
Groups	Plant	Plant	Plant	Avg.	%	Plant	Plant	Plant	Avg.	%
	No. 1	No. 2	No. 3	No.	Difference	No. 1	No. 2	No. 3	No.	Difference
HreX	5	7	8	6.67a	93.37	41	22	20	27.67a	76.49
Buffer Control	107	99	96	100.67b	na	124	106	123	117.67b	na

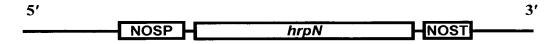
Transformation of Plants and Plant Seeds with a DNA Molecule Encoding a Hypersensitive Response Elicitor Protein

25. In order to investigate whether transforming a plant or plant seed with a DNA molecule encoding a hypersensitive response elicitors results in specific plant

responses, several transformation constructs containing the *hrpN* gene from *Erwinia amylovora* were generated, as described in paragraphs 26-27 below.

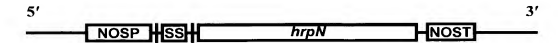
26. A first *hrpN* transformation construct was assembled to include the open reading frame from of the *hrpN* gene inserted behind a nopaline synthase (NOS) promoter, designated NOSP in Figure 1 below, and immediately in front of a NOS terminator, designated NOST in Figure 1 below. The NOS promoter is considered a weak constitutive promoter and has been previously identified. *See* Koncz et al., "The Opine Synthase Genes Carried by Ti Plasmids Contain All Signals Necessary for Expression in Plants," *EMBO J.* 2(9):1597-1603 (1983) (attached hereto as **Exhibit 17**).

Figure 1. Schematic of NOSP-hrpN-NOST Transformation Construct.



27. A second *hrpN* transformation construct was assembled that differed from the construct described in paragraph 25 by the insertion of a tobacco *pr1b* signal sequence, designated SS in Figure 2, between the NOS promoter and *hrpN* open reading frame. The *pr1b* signal sequence has been previously identified. *See* Lund & Dunsmuir, "A Plant Signal Sequence Enhances the Secretion of Bacterial ChiA in Transgenic Tobacco," *Plant Mol. Biol.* 18:47-53 (1992) (attached hereto as **Exhibit 18**).

Figure 2. Schematic of NOSP-SS-hrpN-NOST Transformation Construct.



Experimental Evidence Showing Disease Resistance In hrpN Transgenic Plants

28. As demonstrated by the following experimental evidence in paragraphs 29-30 below, plants grown from seeds harvested from plants transformed with a DNA molecule encoding the hypersensitive response elicitor HrpN from *Erwinia amylovora* exhibited enhanced disease resistance.

29. Arabidopsis Col-0 was transformed with the transformation constructs described above. The constructs were transformed with standard procedures utilizing Agrobacterium transfection. Plants designated 58a8, 58a10, and 58a21 were transformed with the construct described in paragraph 26. Plants designated 60a9, 60a16, and 60a17 were transformed with the construct described in paragraph 27. High hrpN expression transgenic lines were selected by Northern analysis. The lines were confirmed to be homozygous by selection on kanamycin. Prior to initiation of the growth assays, the seeds of each transgenic line and the wild type Arabidopsis were sterilized and subjected to a vernalization treatment in which the seeds were placed at 4°C for approximately four days. All plants were maintained in identical conditions: 16 hours daylight period, 23°C (day)/ 20°C (night), and 50% humidity. Approximately four week after sowing, plants were infiltrated with 10⁶ cfu/ml of Pseudomonas syringae pv. tomato DC3000. Four to six days after inoculation, bacterial concentration were calculated by harvesting 1 cm² of infected leaf tissue, macerating the tissue in 10 mM MgCl, and dilution plating the cell/leaf suspension on King's B plates. Bacterial concentrations in wild type and transgenic lines are shown in Figure 3 below. The data in Figure 3 represents the average of three plants per line and six leaves per plant. Disease proliferation was approximately 70% lower in hrpN transgenic plant compared to non-transgenic wild type plants.

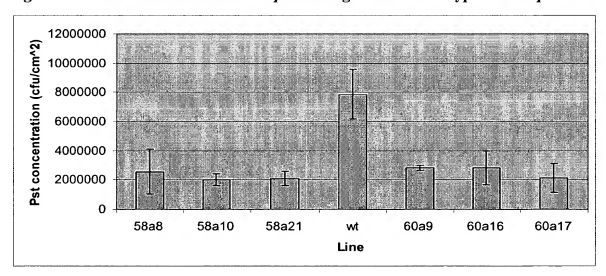


Figure 3. Disease Resistance in hrpN Transgenic vs. Wild Type Arabidopsis

30. Tobacco (Xanthi NN) was transformed with the transformation constructs described above. The constructs were transformed with standard procedures

utilizing *Agrobacterium* transfection. Plants designated 58x2 were transformed with the construct described in paragraph 27. All seeds and plants were maintained in identical conditions: 12 hours daylight period, 26 °C (day)/ 28°C (night), and 50% humidity. Plants were inoculated with TMV as follows. Four leaves per plant were lightly dusted with diatomaceous earth. 100 µl of a 0.42 µg/ml solution of tobacco mosaic virus ("TMV") was applied to the each dusted leaf. The TMV was gently and evenly spread across the leaves. Approximately 5 minutes after inoculation, the plants were lightly rinsed to remove the diatomaceous earth. The number of TMV lesions on the treated leaves was recorded five days after inoculation and is shown in Figure 4 below. *hrpN* transgenic plants had approximately 35% fewer TMV lesions than non-transgenic plant.

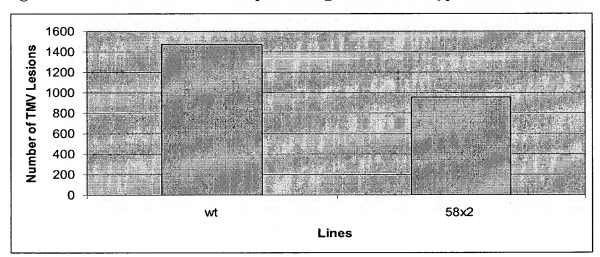


Figure 4. TMV Resistance in hrpN Transgenic vs. Wild Type Tobacco

31. Because disease resistance has been demonstrated for topical application of HrpN of Erwinia amylovora, HrpZ of Pseudomonas syringae, and HreX of Xanthomonas campestris (see supra at ¶¶ 18-20 and 22-24), and transgenic expression of hrpN of Erwinia amylovora and hrpZ of Pseudomonas syringae (see supra at ¶¶ 21 and 28-30), one of ordinary skill in the art would expect other members of this art-recognized class to likewise induce disease resistance in plants following topical application or transgenic expression thereof.

32. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Date:

Zhong-Min Wei

R786765.1

Suresh Gopalan and Sheng Yang He. MSU-DOE Plant Research Laboratory, Michigan State University, East Lansing

Bacterial Genes Involved in the licitation of Hypersensitive Response and Pathogenesis

Intensive molecular genetic studies undertaken in the past 10 years have started to solve many of the puzzles in the area of compatibility and incompatibility between plants and bacterial pathogens. These studies have provided answers to some of the most fundamental questions in plant pathology. What bacterial genes are involved in the establishment of compatibility or incompatibility between plants and necrogenic bacteria? What traits distinguish plant-pathogenic bacteria from saprophysic bacteria? Are these genes and traits common in seemingly very diverse groups of plant-pathogenic bacteria, from soft-rot erwinias to local lesion-forming pseudomonads? In this article, we will discuss some recent advances in understanding the compatibility or incompatibility between plants and necrogenic bacteria (bacteria that cause tissue necrosis). The potential application of these advances to disease management will be addressed briefly. Interested readers should consult other recent reviews (6,8,45,50) for a more technical discussion on this topic.

Plant-Bacteria Interactions: incompatible vs. Compatible

Plant-pathogenic bacteria cause devastating diseases on many important crop plants. Some bacteria, such as Agrobactarium tumefaciens, cause tissue deformation (tumors) by altering hormone balance in infected plant tissues. Other bacteria cause wilt or soft rot by interfering with the function of the plant vascular system or by disintegrating plant tissues, respectively, Many pathovars of Pseudomonas syringae and Xanthomonas compestris cause local lesions on various plant tissues. Disease symptoms caused by most plant-pathogenic baccaria involve plant call death. In this article, only necrogenic bacteria will be

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discussed. Therefore, gall-forming numefaciens and other bacteria that do not cause necrosis will not be addressed.

Plant bacteria interactions can be generally classified as compatible or income patible interactions. In a compatible interaction, a susceptible host plant is infected by a virulent (or compatible) bacterium. resulting in the muitiplication and spread of the besterium in infected plant tissues and the appearance of disease symptoms. In an incompatible interaction, an avirulent (or incompatible) bacterium attempts to infect a resistant host plant or a nonhost plant, but the multiplication and spread of the bacterium are severely restricted. A hallmark of many incompatible interactions is the occurrence of rapid plant cell death ar or near the attempted infection site, a phenomenon known as the hypersenshive response (HR; 16,29). That is, although an. avirulent bacterium is unable to cause typical spreading disease symptoms in a resistant host or nonhost plant, it is able to elicit localized plant cell death. The HR is associated with a wide array of defense responses that may inhibit further pathogen invation, including synthesis of antimicrobial compounds, induction of plant defense genes, and strengthening of the plant cell wall by rapid cross-linking of cell will components (10,32).

Although a true plant-pathogenic bacterium can elicit a dramatic plant responseeither disease or resistance—in a healthy plant with the appropriate genetic background, saprophytic bacteria or bacteria that are pathogenic on organisms other than higher plants are generally unable to initiate any interactions in plants. Of 1,600 known species of bacteria, only about 80 species have been found to cause plant disease (1). What are the features that distinguish with proopents becter's from other Special Taxonomic differences give on clie to the differences in pathugerocky. For example, Erwinia amyioward, the bacterium that causes fire blight, is texonomically more closely related to the human paragens Escherichia coli and Personal specification of another common plant perform f. Appresus.

Genes Controlling Compatibility Between Plants and Bacteria

In the early 1980s, a number of researchers started to: use transposon-mediared mutagenesis, a technique developed in the study of E. coll, to reveal bacterial genes that play important roles in various plant-bacteria interactions. A transposora is a mobile DNA element that can hop in and out of the bacterial chromosome. When a transposon hops into a gene on the chromosome, the gene is physically disrupted and cannot produce a functional product (Fig. 1). If the gene happens to be important in plans-bacterial interactions, the mutant bacterium carrying the disrupted gene will show a defect in initiating normal plant-bacterial interactions.

Using such a mutagenesis technique, Niepoid et al. (35) and Lindgren et al. (33) identified clusters of bacterial genes, known as hrp (for HR and pathogenicity) genes, in the bean pathogens Pseudomonas syringae pv. syringae and P. s. pv. phaseo-: licola respectively. Transposon-induced mutations in hip genes were found to abolish the ability of P. springes to elicit the HR in nonhost plants or to cause disease in host plants (33,35), hip mutants behave very much like bacteria that have no apparent interactions with plants, such as E. coli. The identification of hip genes suggested that the molecular mechanism(s) underlying bacterial pathogenicity and bacterial elicitation of plant disease resistance may involve the same bacterial

hap genes have been isolated from many plant-pathogenic bacteria and have been characterized most extensively from P. s. pv. syringae, P. z. pv. phaseolicola, Pseucomonas solanacearian (which causes wilt : in many solanaceous plants), Xanthomonas campenris pv. vesicatoria (which causes bacterial spot on tomato and pepper), and E. anylovora (6,8,45). Isolation (cloning) of hop genes was accomplished by inserting random genomic DNA fragments from a wild-type, plant-pathogenic bacterium into a cloning vector, followed by introduction of cloned DNA fragments into thrp mutants

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(Fig. 1). If a cloned DNA fragment carries a wild-type copy of the mutated harp gene in an hip mutant, it will produce a functional hrp gene product and therefore complement the mutated hrp gene located in the chromosome (Fig. 1). Surprisingly, the cloned hep clusters from P. 1 pv. syringae 61 and E amylovora 321 enabled nonpathogens (e.g., E coll or Pseudomonas fluorescens) to elicit the HR in plants (5.24). The functional cloning of these two hep clusters in E coli revealed that the minimum number of genes required for elicitation of the HR by plant-pathogenic bacteria is carried on a DNA fragment about 25 to 30 kb in length, a very small portion of the bacterial genome (which is normally about 4,000 to 5,000 kb).

DNA-DNA hybridization studies indicate that at least some hop genes are similar among necrogenic bacteria belonging to different genera (P. syringat, E. amylovora, Erwinia siewartii, P. solanocearu and X campestris) (31). Recent DNA sequence studies confirm that many hap genes cloned from diverse plant-pathogenic bacteria are homologous (23,46). Thus, hip genes appear to be universal among diverse necrosis-causing gramnegative bacterial pathogens of plants.

Biochemical Functions. of hrp:Genes

The biochemical functions of kry genes have remained a puzzle until recently. DNA sequencing has played a major role. in the determination of many lup sens functions. As will be discussed, many top genes have striking similarities with ge of known function. Figure 2 shows the gene organization and likely functions of hop genes of P. + pv. syringae (23). There are at least 25 Jop genes in this bacterium. Based on DNA sequence similarity to other known genes and subsequent biochemical and molecular characterization, we now know that Jup genes have at least three biochemical functions: gene regulation, protein secretion, and production of HR

elicitor proteins. 1. Gene regulation. It was discovered that hrp genes either are not expressed or are expressed at very low levels (i.e., they make very low levels of protein products) when bacteria were grown in nutrient tich bacteriological media, whereas they are highly expressed when bacteria are in the intercellular space (apoplast) of plant dissues (25,37,41,46,48,52,53). What conditions in plant tissues induce the expression of hip genes, and how do bacteria detect these inducing conditions? Unlike viruses, nematodes, and many fungi, plant-pathogenic bacteria do not invade living plat: cells. Therefore, signal exchanges between plant cells and bacteria must occur in (or through) the apoplast outside the plant cell. A number of laboratories have observed that induction of P. syringae hrp genes and he achieved by using artificial

minimal media lacking complex nitrogen nutrients, indicating that lack of nutrients in the plant apoplast may be the signal for the induction of hrp. genes (25,37,52,53). Specific compounds (e.g., sucrose and sulfur-containing amino acids) present in the plant apoptast may also serve as signals for the induction of X c. pv. vesicotoria hrp genes (41). The induction of hrp genes in the nutrient-poor plant apoplast or in artificial minimal media indicates that hep genes may be involved in bacterial nutrition in planta.

How do bacteria sense the plant apoplast environment? It was found that at least three of the 25 hip gene products are involved in detection of the apoplast environment by P. syringae, hrpl., hrps, and hrpR (18,51; Fig. 2). The hrpS and hrpR are among the first two hop genes to be expressed once bacteria enter plant tissues (51,52). It has been bypothesized that the HirpS and HirpR proteins, once produced, bind to the "promous" sequence of the hirpL gene to "promote" the production of the HrpL protein (51). Once the HrpL protein is produced, it activates promoters of other hop genea and some bacrecial avirulence (avr; genes, which determine gene-for-gene interactions between bacteris and plants (25.26.38,40,51; Fig. 3). Not all bacterial ave genes are regulated by hirp genes (30). Interestingly, hrpS and hrpR

are similar in sequence to a family of bacterial proteins that regulate genes involved in diverse metabolic functions, including genes involved in nutrient transport and metabolism (18,51). The sequence similarity of hrpS and hrpR with gene regulators involved in nutrition appears to support the hypothesis that hip genes are involved in bacterial nutrition in the mutrient-poor plant apoplast. This hypothesis is further supported by the observation that the expression of hop genes can be turned off by complex mitrogen sources, tricarboxylic sold cycle intermediates, high osmolarity, and neutral pH, some of which are characteristic of rich bacterial media (25,37,41,46,52,53).

An hrpS homolog has been found in a very different bacterium, E. amylovora (S. V. Beer, personal communication). In P. solanocearum, a different hrp gene (hrpB). was found to be involved in the detection of the plant apoplast (15). Thus, different. bacteria may or may not use the same mechanism to detect the apparently similar environment in the plant apoplast.

2. Protein secretion. One surprising finding from the sequence analysis of hrp genes was that many hop genes show striking similarities to those involved in the secretion of proteinsceous virulence factors in human and animal pathogenic bacteria (12,17,22,39,46). Most plant-pathogenic

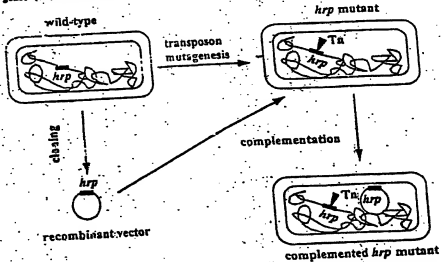


Fig. 1. Disgram of molecular techniques commonly used in the cioning of hrp genes. A wild-type bacterium is mutagenized by random insertion of a transposen (Tn) into its genome. When a transposon inserts into 1 wild-type hrp gene (in red), it physically disrupts the hip gene (in green). The transposon-inserted hip gene cannot produce & disrupts the hip game (in green). The wansposin-inserted hip game cannot produce a functional product, and the resulting backerium is called a hip mutant. The hip mutant functional product, and the resulting backerium is called a hip mutant. The hip mutant functional product and hip mutant for cause can no longer induce the hypersolative response (HR), in resistant plants or cause can no longer induce the hypersensitive response (HR) in resistant plants or cause disease in susceptible plants. It is losted (pione) the hrp gene identified by transposen numbers of gene jibrary is established by inserting places of the wild-type generally. Into a cloning vector (indicated by a circle). The vector parrying to reign generally is then introduced into the hrp mutant. If a premishent vector happens to carry a wild-type copy of the mutant hrp gene, it will represent a functional hrp gene product lacking in the hrp mutant, thus recovering the patient of the mutant to induce the HR in resistant plants and to cause disease in ability of the mutant to induce the HR in resistant plants and to cause disease in susceptible plents. The hrp mutant phenotype is therefore complemented by this recombined vector.

bacteria that cause necrosis are gramnegative, that is, they have two cell membranes enveloping the cytoplasm. For any large molecule (e.g., a protein) to go through a lipid membrane, a special secretion apparatus or channel composed of many proteins must be assembled across both cell membranes. Gram-negative plant pathogenic bacteria are known to make several types of secretion apparatus, For example, Erwinia chrysanthemi, a tocte rium that causes soft rot, makes one type of secretion apparatus for proteases and another for plant cell wall-degrading enzymes (21,39). Both types of secretion apparatus are widely conserved among many other bacteria, including human pathogens such as E coli and Pseudomonas aeruginasa (21,39). The hrp genes were found to specify a third type of secretion apparatus, the Hrp secretion apparams, which appears to be similar to the one discovered in several human-pathogenie bacteria, including Yersinia spo-(12,17,22,39,46). Interestingly, although the regulatory hrp genes in different bacteria may be different (hrpS, hrpR, and hrpl in P. syringae versus hrpB in P. solanacearum), most hrp genes involved in the assembly of the Hip secretion apparates are similar among diverse plant-pathogenic bacteria. This suggests that although different bacteria may detect the plant apoplast environment in their own unique ways, they nevertheless produce similar types of protein secretion apparants.

3. Production of elicitor proteins. The T discovery of the novel Hrp secretion appa-

ratus raised an immediate question: What are the proteins that traverse it? Since hap genes are essential for bacteria, both to elicit the plant HR and to cause disease, it was expected that some of the proteins that traverse the Hrp secretion apparatus may be elicitors of plant HR and that others may be involved in causing necrosis during pathogenesis. Wei et al. (47) first provided evidence that one of the E amylovorp hap genes (hrp.V) encodes a proteinaceous elizator (harpin). Harpin elicita MR accrusio when injusted into the apoplast of appropriate plasss (47). Although no hrpN gene homolog could be found in P. syringer, ... another approteinaceous a HR elicitor (harpings) was identified and was shown to be encoded by a different hop gene, ArpZ (20.36). Furthermore, harpines was the first extracellular protein shown to be recreiment is the Hop secretion apparatus 120% Athirduscerial protein elicitor of the HR was identified in P. solanacearum and was mailed PopA1-(2). The E. amylovoro harpin, P. 1. py, syringae 61 harpings, and P. solariacearum PopAI, although largely dissimilar in primary sequences, share similar properties that may be important in their HR elicitor activities. They are all heat stable, glycine rich, and hydrophilic. Homology of E. amylovera harpin and P. z. pr.: . wringer .. 61 harping have been identified in other pathogenic strains that belong to the genus Erwinia and the species ? syringae, respectively (4,20). Titus, at least three proteins that traverse the riry secretion apparatus of three diverse bacteria elicit the HR. SEPT T

The Search for Proteins that Traverse the Hrp Apparatus

As mentioned earlier, bacterial mutants defective in the Hrp secretion apparatus are unable to elicit the HR in resistant plants and to cause disease in susceptible plants. The question is, how many proteins are secreted via the Hrp secretion apparants? If harpins and PopA are the only proteins that traverse the Hrp secretion apparatus, then morations in the genes that make harpins and PonA would also eliminate the ability of bacteria to elicit the HR in resistant plants and to cause disease in host plants. However, if there are other pathogenicityor HR-related proteins secreted via the Hrp apparatus, mutations in only harpin- or PopA-encoding genes would not completely abolish the ability of bacteria to elicit the HR in resistant plants or to cause disease in host plants. Wei et al. (47) reported that mutations in the gene coding for harpin of E. amylovora destroyed the ability of the bacteria both to trigger the HR in resistant monhost tobacco and to cause disease in susceptible pear fruits. Mutations in the gene coding for harpinges of E. chrysanthemi prevented the bacterium from triggering the HR in the nonhost tobacco and reduced the ability of the bacterium to initiate disease lesions in host plants (4). In the case of barpiness of P. syringae, mutation analysis has been complicated by the complex gene structure and organization surrounding the hrpZ gene. Conclusive data regarding the role of : harpings in plant-P. syringes interactions are yet to be shown. PopAl was shown to

Pseudomonas syringae hrp gene cluster

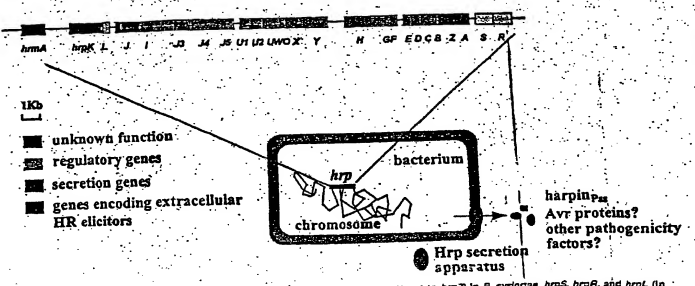


Fig. 2. hrp genes of Pseudomonas syringse. There are at least 25 hrp genes (hrpA to hrpZ) in P. syringse, hrpS, hrpR, and hrpL (in yellow) are involved in the detection of the plant acceptant environment and in the activation of all other hrp genes, and possibly other pathogenicity-related genes. Most other hrp genes (in red) are involved in the assembly of the hrp secretion apparatus in the bacterial envelope, through which travels a newly discovered class of bacterial virulence/avirulence proteins (in green), including hrpZ.

dispensable for pathogenicity of P. solanacearum in the susceptible host plant, tomato, or for HR elicitation in the nonhost plant, tobacco (2), indicating that there must be other HR-elicitors and pathogenicity factors that traverse the Hrp secretion apparatus in this bacterium. Further examination indicated that PopA1 may function as an avirulence gene, medisting gene-for-gene interaction in certain Penunia P. solonoceanon interactions (2.45). Thus, the Hrp secretion apparatus in each bacterium may secrete a different number of proseins. Identification of other proteins that traverse the Hrp secretion apparatus is now an active research area and is essential for a complete understanding of hrp-mediated plant-bacterial interactions.

Factors Modifying hrp Gene-Mediated Compatibility

Two broad classes of bacterial genes may superimpose their functions on the hrp gene-mediated compatibility or incompatibility between plants and bacterist avr genes and various virulence genes. The ave genes mediate genotype-specific incompatibility in resistant bost plants. Virulence genes promote the production of disease symptoms and are usually needed for the full virulence of bacteria.

Bacterial avr. Genes.

Flor (14) formulated the gene-for-gene hypothesis in his work on flax-flax rust interactions. Flor hypothesized that the resistance of a given flax cultivar to a given fungal race is the result of the interaction between a fungal ovr gene and a corresponding flax resistance gene. Therefore, the interactions between the plant's. resistance genes and the pathogen's aur genes would limit the host range of the pathogen. Staskawicz et al. (44) first cloned an avr gene from a soybean becterial pathogen, Pseudomonas syringas pv. glycinea, and showed that the cloned aver gene could convert virulent. P. 2. pv. glycinea strains that cause disease into svirulent strains that elicit the HR in certain soybean cultivars carrying the corresponding resistance genes, thus validating the role of our genes in controlling bost range. Since then, numerous our genes have been cloned from plant-pathogenic bacteria (27). Several plant resistance genes have also been cloned using molecular genetic approaches (e.g., 34,43).

What is the relationship between the cor genes and hip genes, both of which are involved in eliciting the HR? Several laboratories have observed that avr genes cannot trigger the genotype-specific HR in hrp mutants, i.e., avr genes depend on functional hrp genes for expressing their phenotype (25,26,28,38,40). There are several ways of explaining such dependence (Fig. 4). One possibility is that Avr proteins are dependent on the Hrp secretion apparatus for secretion. Alternatively, Avr function requires a prior plant response

elicited by the hyp-controlled extraceliular factors (such as harpins). A third possibillity is that Avr proteins, with no HReliciting activity by themselves, cause the cultivar-specific HR by, either covalently modifying harpins or modulating the expression of harpins in a plant resistance gene-dependent manner yet to be understood. Finally, it is also possible that Ave proteins are secreted directly into the plant cell with the help of harpins, assuming that receptors for Avr proteins are inside the plant cell. Studies are being carried out to resolve these possibilities.

Bacterial Virulence Factors

The genetic diversity of plant-patho-genic bacteria is reflected in their ability to cause diverse disease symptoms ranging from soft not to tissue necrosis to wildfire." These diverse disease symptoms are likely the result of the action of several, sometimes unique, virulence fac-tors produced by a given bacterium in addition to kep-controlled pathogenicity

factors. For example, research from many laboratories has shown that toxin production plays an important role in the formation of chlorosis and necrosis (3,19,49). Extracellular polysaccharides may be involved in the formation of water-scaking lesions (11,13) and in the production of will symptoms by clogging the plant vascular system (9). Plant cell wall-degrading enzymes are responsible for tissue disintegration and the appearance of the soft-rot symptom (7). Plant bormones produced by plant-pathogenic bacteria are involved in the induction of tissue deformation (42).

Both hip genes and batterial virulence factors are necessary for disease symptom production, but what is the relationship between them? A logical relationship would be that hrp-controlled extracellular factors are involved in obtaining mutrients in early stages of pathogenesis, whereas other virulence factors drive the initial compatible stage into a fully compatible one, leading to the production of various disease symptoms. At least two lines of

. plant apoplast signais

plant apoplast

bacterial cell wall bacterial cytoplasm

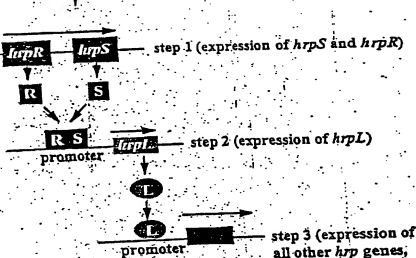


Fig. 3. Diagram of the signal transduction cascade in the detection of the plant apoplast environment by Pseudomones syringes. The plant apoplast environment (limited nutrients and/or cartain unique compounds) activates the expression of hrps (step 2). The hrpL gene product (L), in turn, binds to promoters of other hrp genes, syr gener, and other becturius pathogenicity-related genes to promote the expression of general send point pectation paragraphically pectage plant-bectaria interactions (stap 3). these general resulting in the initiation of diverse plant-bectaria interactions (stap 3). Madified from Xiao et al. (51).

avr. genes, and other

genes)

pathogenicity-related

evidence seem to support this relationship. First, hrp genes are highly conserved among diverse plant-pathogenic bacteria, whereas virulence factors vary greatly among bacteria. Second, while mutations in the hrp gene completely abolish both bacterial pathogenicity and elicitation of the HR, mutations in virulence genes (e.g., mxin-production genes) often do not eliminate pathogenicity and have no effect on bacterial elicitation of the HR (3,19,49).

hrp Gene Functions and Disease Management

A major reason for discovering bacterial and plant factors critical for bacterial pathogenesis and plant resistance is to develop novel and environmentally safe strategies for controlling plant diseases. The discovery that the Hrp secretion apparabus is crucial to bacterial pathogenesis provides a foundation for designing novel. chemicals and antibodies that would block

the assembly of the Hrp secretion apparatus or the passage of bacterial virulence proteins through it. Alternatively, susceptible crop plants could be genetically engineered with genes encoding proteinaceous HR elicitors, such as harpins, under the control of plant promoters inducible by virulent pathogens. If this approach were successful, the HR or resistance would be triggered in otherwise compatible interactions, limiting disease development.

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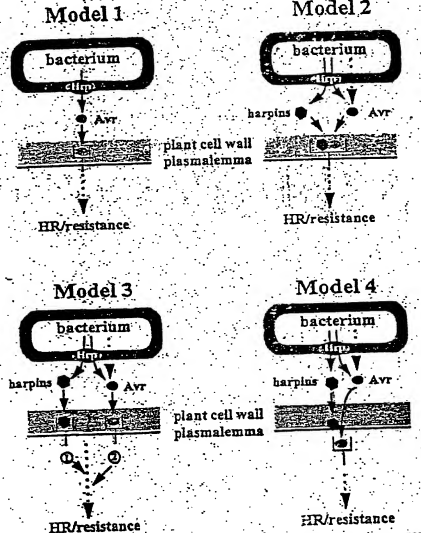


Fig. 4. Worlding models for possible interactions between top genes and sur genes. Model 1: Avr signals (Avr proteins or their enzymatic products) are secreted through . the Hrp secretion apparatus to elicit the hypersensitive response (HR) and resistance. Model 2: Harpins and Avr signals modify each other before interacting with plant receppors. Avr. signals may or may not be recreted to the Hrp secretion apparatue. Model 3: Harpine and Avr signals interest, with respective plant receptors. Plant response elicited by harpins must precede plant response elicited by Avr. signals. Avr. signals may or may not be secreted via the Hrp secretion apparatus. Model 4: Avr. proteins are secreted into the plant call with the help of harpins. Avr signals may or may not be secreted via the Hirp secretion apparatus. In models 1 to 3, receptors for Avr proteins are presumed to be on the plant cell surface. In model 4, receptors for Avr proteins are inside the plant cell.

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Erwinia chrysanthemi Harpin_{Ech}: An Elicitor of the Hypersensitive Response that Contributes to Soft-Rot Pathogenesis

David W. Bauer, Zhong-Min Wei, Steven V. Beer, and Alan Collmer

Department of Plant Pathology, Cornell University, Ithaca, NY 14853-4203 U.S.A. Received 14 December 1994. Accepted 8 March 1995.

Mutants of the soft-rot pathogen Erwinia chrysanthemi EC16 that are deficient in the production of the pectate lyase isozymes PelABCE can elicit the hypersensitive response (HR) in tobacco leaves. The hrpN_{Ech} gene was identified in a collection of cosmids carrying E. chrysanthemi hrp genes by its hybridization with the Erwinia amylovora hrpNEs gene. hrpNEch appears to be in a monocistronic operon, and it encodes a predicted protein of 340 amino acids that is glycine-rich, lacking in cysteine, and highly similar to HrpN₅₀ in its C-terminal half. Escherichia coli DH5a cells expressing hrpN_{Ech} from the lac promoter of pBluescript II accumulated HrpNeca in inclusion bodies. The protein was readily purified from cell lysates carrying these inclusion bodies by solubilization in 4.5 M guanidine-HCl and reprecipitation upon dialysis against dilute buffer. HrpN $_{Ech}$ suspensions elicited a typical HR in tobacco leaves, and elicitor activity was heat-stable. Tn5-gusAI mutations were introduced into the cloned $hrpN_{Ech}$ and then marker-exchanged into the genomes of E. chrysanthemi strains AC4150 (wild type), CUCPB5006 (ΔpelABCE), and CUCPB5030 (ΔpelABCE outD::TaphoA). hrpNEch::Ta5-gusA1 mutations in CUCPB5006 abolished the ability of the bacterium to elicit the HR in tobacco leaves unless complemented with an hrpN_{Ech} subclone. An hrpN_{Ech}:: Tn5-gusAl mutation also reduced the ability of AC4150 to incite infections in witloof chicory leaves, but it did not reduce the size of lesions that did develop. Purified $HrpN_{Ech}$ and E. chrysanthemi strains CUCPB5006 and CUCPB5030 elicited HR-like necrosis in leaves of tomato, pepper, African violet, petunia, and pelargonium, whereas $hrpN_{Ech}$ mutants did not. HrpNeck thus appears to be the only HR elicitor produced by E. chrysanthemi EC16, and it contributes to the pathogenicity of the bacterium in witloof chicory.

The hypersensitive response (HR) is a rapid, localized necrosis that is associated with the active defense of plants against many pathogens (Kiraly 1980; Klement 1982). The HR elicited by bacteria is readily observed as a tissue collapse if high concentrations (≥10⁷ cells per milliliter) of a limited-host-range pathogen like *Pseudomonas syringae* or *Erwinia amylovora* are infiltrated into leaves of nonhost plants (ne-

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crosis occurs only in isolated plant cells at lower levels of inoculum) (Klement 1963; Klement et al. 1964; Turner and Novacky 1974; Klement 1982). The capacities to elicit the HR in a nonhost and to be pathogenic in a host appear linked. As noted by Klement (1982), these pathogens also cause physiologically similar, albeit delayed, necroses in their interactions with compatible hosts. Furthermore, the ability to produce the HR or pathogenesis is dependent on a common set of genes, denoted hrp (Lindgren et al. 1986; Willis et al. 1991). Consequently, the HR may hold clues to both the nature of plant defense and the basis for bacterial pathogenicity.

The hrp genes are widespread in gram-negative plant pathogens, where they are clustered, conserved, and in some cases interchangeable (Willis et al. 1991; Bonas 1994). Several hrp genes encode components of a protein secretion pathway similar to one used by Yersinia, Shigella, and Salmonella spp. to secrete proteins essential in animal diseases (Van Gijsegem et al. 1993). In E. amylovora, P. syringae, and P. solanacearum, hrp genes have been shown to control the production and secretion of glycine-rich protein elicitors of the HR (He et al. 1993; Wei and Beer 1993; Arlat et al. 1994).

The first of these proteins was discovered in E. amylovora Ea321, a bacterium that causes fire blight of rosaceous plants, and was designated harpin (Wei et al. 1992). Mutations in the encoding hrpN gene revealed that harpin is required for E. amylovora to elicit the HR in nonhost tobacco leaves and incite disease symptoms in highly susceptible pear fruit. The P. solanacearum GMI1000 PopA1 protein has similar physical properties and also elicits the HR in leaves of tobacco, which is not a host of that strain (Arlat et al. 1994). However, P. solanacearum popA mutants still elicit the HR in tobacco and incite disease in tomato. Thus, the role of these glycine-rich HR elicitors can vary widely among gram-negative plant pathogens.

E. chrysanthemi is unlike the bacterial pathogens that typically elicit the HR because it has a wide host range, rapidly kills and macerates host tissues, and secretes several isozymes of the macerating enzyme pectate lyase (Pel) (Barras et al. 1994). Nevertheless, PelABCE and Out (pectic enzyme secretion pathway) mutants of E. chrysanthemi EC16 cause a typical HR (Bauer et al. 1994). Furthermore, elicitation of the HR by E. chrysanthemi is dependent on an hrp gene that is conserved in E. amylovora and P. syringae and functions in the secretion of the E. amylovora harpin (Wei and Beer 1993; Bauer et al. 1994). Mutation of this gene significantly reduces the ability of E. chrysanthemi to incite lesions in susceptible

witloof chicory leaves. These observations suggest that E. chrysanthemi also produces a harpin. We report here the cloning, characterization, and mutagenesis of the E. chrysanthemi $hrpN_{Ech}$ gene and an investigation of the role of its product in plant interactions.

RESULTS

Molecular cloning of the E. chrysanthemi hrpN_{Ech} gene.

We previously isolated 18 cosmids containing E. chrysan-themi DNA sequences hybridizing with the E. amylovora hrp cluster (Bauer et al. 1994). The pattern of restriction fragments released from these cosmids indicated they all contained overlapping inserts from the same region of the E. chrysan-themi genome (data not shown). The cosmids were probed in colony blots with a 1.3-kb HindIII fragment from pCPP1084.

8 0 1 T 1 K A H 1 G G D L 61 TCTCCCGGTCTGCCGCTGCTCAGGGACTCAAAGGACTGAATTCCCGCGCTTCAT V S G L G L G A Q G L K G L N S A A S 121 TCGCTTCCAGCGTGCATAAACTGAGCAGCACTGCATAAGTTCACCTCCGCGCTCA L G S S V D K L S S T I D K L T S A L 181 CGATGATGTTTCCCCGCGCGCGCGCGCGCGCGCGCGCCCCACCTCCAAGGCGCGTCC S M N F G G A L A Q G L G A S S K G L 241 TCAGCAATCAACTGCCCAGTCTTTCCCCAATGCCGCCCACCCTGCCACCACCTCCCAACCTCCC N S N Q L G Q S F G N G A Q G A S N L 1	CGC S
V S G L G L G A Q G L K G L N S A A S 121 TCGCTTCCAGCCTGGATAAACTGAGCAGCACCATCGATAAGTTGAGCTCGGCGCTGAL L G S S V D K L S S T I D K L T S A L T 181 CGATGATGTTTGCGCGCGGCGCTGGGGGGGGGGGGGGGG	S TT T CA
121 TCCCTTCCACCCTCGATAAACTGACACACCATCGATAAGTTGACCTCCGCCCTCAL L C S S V D K L S S T I D K L T S A L 181 CGATGATGTTTCCCCCGCGCGCTCCCCGAGGCGCTGGGCCCCAGCCTCCAAGGCCCTCCC S H N F G G A L A Q G L C A S S K G L C 241 TGAGGAATCAACTGGGCGAGTCTTTCCGCAATGGCGCCCAGGGTGCGAGGAACCTTCCC	TTI T
L G S S V D K L S S T I D K L T S A L T 181 CGATGATGTTTGGGGGGGGGGGGGGGGGGGGGGGGGGGG	CA
L G S S V D K L S S T I D K L T S A L T 181 CGATGATGTTTGGGGGGGGGGGGGGGGGGGGGGGGGGGG	CA
181 CGATGATGTTTCGCCGCGCGCGCGCGCGGGGGGGGGGGG	CA
S M M F C G A L A Q C L C A S S K C L (241 TGAGGAATGAAGGGGGGGGGGGGGGGGGGGGGGGGGGG	CA ;
S M M F C G A L A Q C L C A S S K C L (241 TGAGGAATGAAGGGGGGGGGGGGGGGGGGGGGGGGGGG	CA ;
241 TCAGCAATCAACTGGGGGGGGGGGGGGGGGGGGGGGGGG	;
241 TCAGCAATCAACTCGCCCAGTCTTTTCGCCAATGCCGCCCAGGCTGCCAGCAACCTGCC	
M S N O L C O S F G N C A O C A C N	
	ΑI
	•
301 CCCTACCGAAATCCCCCGCGGCGATGCGTTGTCAAAAATGTTTGATAAAGCCCTGGACG	Tr
S V P K S G G D A L S K M F D K A L D I	
361 TECTOCCTCATGACACCCTGACCAAGCTGACTAACCAGAGCAACCAAC	*
LLGHDIVIKLINQSNQLANS	
4> 439; :Tn5-gugAI	
421 TGCTGAACGCCAGCCAGATCACCCAGGCTAATATGAATGCGTTCGGCAGCGCTGTGAA	CA
M L N A S Q M T Q G N H N A F G S G V H	
481 ACGCACTCTCCTCCATTCTCGGCAACGGTCTCGGCCACTCCATCAGTCGCTTCTCTCA	CC
N A L S S I L C N C L C Q S M S C F S Q	
546::Tn5-gusAl <4	
541 CTTCTCTGGGGGCAGGCCGCTTGCAGGCCCCTGAGCGGGGGGGG	CC
P S L C A C C L Q C L S C A C A F N Q L	
601 GTAATGCCATCGGCATCGGCGTCGGGCACAATGCTGCGCTGAGTCCGTTGAGTAACGT	٠.
G N A I G M G V C Q N A A L S A L S N V	
661 GCACCCACGTAGACGGTAACAACCGCCACTTTGTAGATAAAGAAGATCGCGGCATGGC	
5 THVDGNNRHFVDKEDRGHA	
721 AAGAGATGGGCCAGTITATGGATGAGTATGCGGAAATATTGGGTAAACGGGAATAGCA	:A
KEICQFHDQYPEIFGKPEYO	
· ·	
781 AAGATGGCTGGAGTTGGCGGAAGACGGACGACAAATCCTGGGCTAAAGGGCTGAGTAA	ıc
K D G V S S P K T D D K S W A K A L S K	
*** *** · · · · · · · · · · · ·	
841 CCCATCATGACCCTATCACCCGCCCCAGCATGCACAAATTCCGTCAGGCGATGGGTATC	À
841 CGGATGATGACGGTATGACGGGGCCCAGCATGCACAAATTCCGTCAGGGGATGCGTATG	A
PDDDGHTGASHDKFRQAHGH	
P D D D G H T G A S H D K F R Q A H G N	
PDDDGHTGASHDKFRQAHGH	
PDDDGMTGASSMDKFRQANGG 901 TCAAAAGCGCGGGGGGGGGGGGGGGATACCAACCTGAACCTGCGGGGGGGG	c
P D D D G M I G A S M D K F R Q A M G M 901 TCAAAAGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	c
PDDDGMTGASSMDKFRQANGG 901 TCAAAAGCGCGGGGGGGGGGGGGGGATACCAACCTGAACCTGCGGGGGGGG	c
901 TCAAAAGCGCGCGCGCGCGCGCGCAATACCCAACCTCGACCCTCGCGCGGGGGGGG	ic ic
901 TCAAAAGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	ic ic
901 TCAAAAGCGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG	ic ic

Fig. 1. DNA sequence of $hrpN_{Ech}$ and predicted amino acid sequence of its product. Underlined are the putative ribosome-binding site, the N-terminal amino acids confirmed by sequencing the product of pCPP2172, and a potential rho-independent transcription terminator. The location and orientation of two Tn5-gusA1 insertions are also indicated and are numbered according to their location in the $hrpN_{Ech}$ open reading frame. The accession number for hrpN is L39897.

which contains the *E. amylovora hrpN* gene (Wei et al. 1992). pCPP2157, one of the three cosmids hybridizing with the probe, was digested with several restriction enzymes, and the location of the hrpN_{Ech} gene in those fragments was determined by probing a Southern blot with the *E. amylovora HindIII* fragment. Two fragments, each containing the entire hrpN_{Ech} gene, were subcloned into different vectors: pCPP2142 contained an 8.3-kb SalI fragment in pUC119 (Vieira and Messing 1987), and pCPP2141 contained a 3.1-kb PsrI fragment in pBluescript II SK(-) (Stratagene, La Jolla, CA).

Sequence of hrpN_{Ech}.

The nucleotide sequence of a 2.4-kb region of pCPP2141 encompassing $hrpN_{Ech}$ was determined. The portion of that sequence extending from the putative ribosome-binding site through the hrpN_{Ech} coding sequence to a putative rho-independent terminator is presented in Figure 1. The typical ribosome-binding site, consisting of GGAAA, was located eight bases upstream of the ATG translational initiation codon. A consensus hrp promoter sequence of GGAACC(N)16CACTCA (Bonas 1994) was found 97 bases upstream of the open reading frame (ORF), suggesting that hrpN_{Ech} is a monocistronic operon. hrpN_{Ech} codes for a predicted protein that has a molecular mass of 34.3 kDa, is rich in glycine (16.2%), and is lacking in cysteine. Comparison of the amino acid sequences of the predicted $hrpN_{Ea}$ and $hrpN_{Ech}$ products revealed extensive similarity, particularly in the C-terminal halves of the proteins (Fig. 2). The overall identity of the

```
Ech ........MQITIKAHIGGDLCVSGLGLGAQGLKGLNSAASSLGSSVDKL 42
E4 MSLNTSCLGASTMQISIGGAGGNNGL.LGTSRQNAGLGGNSALGLGGGNQ 49
Ech SSTIDKLTSALTSMMF......GCALAQGLGAS.SKGLGMSNQLGQSFG 84
   NDTVNQLAGLITGHOODHYSHMGGGGLMGGGGGGCGLGNGLGGSGGLGEGLS 99
..... HDTVTKLTNQSNQLANSHLNAS...... QHTQCNNNAFG 150
   STSDSSDPHQQLLKMFSEIHQSLFCDCQDGTQGSSSCCKQPTEGEQNAYK 199
Ech CACAFNQLCNAIGHGVGQNAALSALSNVSTHVDCNNRHFVDKEDRGNAKE 236
  Ech IGQFNDQYPEIFGKPEYQKDGWSSPKTDDKSWAKALSKPDDDGHTGASHD 286
  Ech KFRQAMGMIKSAVAGDTGNTNLNIRGAGGASLGIDAAVVGDKIANMSLGK 336
  -|--|-|||-::||||||:
QFNKAKGHIKRPMAGDTGNGNLH....
                       ||..|. : . |.
...DAVPVVLRWVLMP... 385
Ech LANA 340
```

Fig. 2. Predicted amino acid sequences of the hrpN products HrpN_{EA} (Ech) of Erwinia chrysanthemi and HrpN_{Ea} (Ea) of E. amylovora, aligned by the Gap program of the Genetics Computer Group Sequence Analysis Software Package (Devereaux et al. 1984). Two dots denote greater similarity than one dot.

hrpN genes and proteins was 66.9 and 45.5%, as determined by the FASTA and Gap algorithms, respectively (Devereaux et al. 1984; Pearson and Lipman 1988).

The direction of hrpN_{Ech} transcription, the size of the predicted product, and the translation start site were confirmed by recloning the 3.1-kb PstI fragment from pCPP2157 and selecting a clone with the fragment in pRluescript II SK(-) in the opposite orientation from pCPP2141, to produce pCPP2172. E. coli DH5α(pCPP2172) expressed hrpN_{Eth} from the vector lac promoter and produced high levels of a protein with an estimated molecular mass of 36 kDa in sodium dodecyl sulfate (SDS) polyacrylamide gels, which is close to the predicted size (Fig. 3). Furthermore, the 10 N-terminal amino acids of the 36-kDa protein, determined by microsequencing following purification as described below, corresponded with the predicted N terminus of HrpN_{Ech}. As expected, no Nterminal signal sequence for targeting to the general export (Sec) pathway was discernible in the HrpN_{Ech} sequence, and our data showed no evidence of processing of the N terminus.

Purification of the $hrpN_{Ech}$ product and demonstration of its HR elicitor activity in tobacco.

When DH5 α (pCPP2172) cells were disrupted by sonication and then centrifuged, most of the HrpN_{Ech} protein sedimented with the cell debris. However, soluble HrpN_{Ech} could be released from this material by treatment with 4.5 M guanidine-HCl. This suggested that the protein formed inclusion bodies which could be exploited for purification. As detailed in Materials and Methods, we found that HrpN_{Ech} reprecipitated when the guanidine-HCl was removed by dialysis against dilute buffer. The HrpN_{Ech} precipitate could be washed and resuspended in buffer, in which it formed a fine suspension. SDS polyacrylamide gel analysis revealed the suspension to be electrophoretically homogeneous HrpN_{Ech} (Fig. 3).

Cell-free lysates from E. coli DH5α(pCPP2172) cells grown in Luria-Bertani medium were infiltrated into tobacco

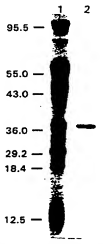


Fig. 3. Sodium dodecyl sulfate (SDS) polyacrylamide gel of purified $\operatorname{HrpN}_{Ech}$. Purified $\operatorname{HrpN}_{Ech}$ was solubilized in SDS loading buffer, electrophoresed through a 12% polyacrylamide gel, and stained with Coomassie Brilliant Blue. Lane 1, molecular weight markers (mid-range markers from Diversified Biotech, Boston, MA), with sizes in kilodaltons shown to the left; lane 2, $\operatorname{HrpN}_{Ech}$.

leaves. Necrosis typical of the HR developed within 18 h, whereas leaf panels infiltrated with identically prepared lysates of DH5α(pBluescript SK-) showed no response (data not shown). The suspension of purified HrpN_{Ech} at a concentration of 336 µg/ml also caused a necrotic response within 18 h that was indistinguishable from that caused by E. chrysanthemi CUCBP5030 or cell-free lysates from E. coli DH5α(pCPP2172) (Fig. 4). Tobacco plants vary in their sensitivity to harpins, and elicitation of the HR by $HrpN_{Ech}$ at lower concentrations was found to be variable. Consequently, a concentration of 336 µg/ml was used in all subsequent experiments. The concentration of $HrpN_{\mathcal{E}_{ch}}$ that is soluble in apoplastic fluids is unknown. To determine the heat stability of HrpN_{Ech}, the suspension of purified protein was incubated at 100° C for 15 min and then infiltrated into a tobacco leaf. There was no apparent diminution in its ability to elicit the HR (data not shown). These observations indicated that HrpN_{Ech} is sufficient to account for the ability of E. chrysanthemi to elicit the HR in tobacco.

$hrpN_{Ech}$ mutants fail to elicit the HR in tobacco.

E. coli DH10B(pCPP2142) was mutagenized with Tn5-gusA1 (Sharma and Signer 1990). Plasmid DNA was isolated

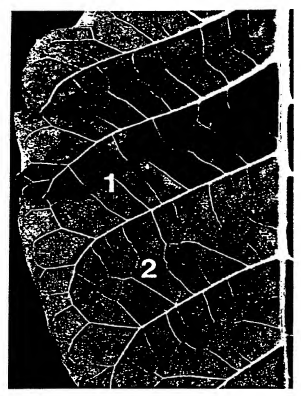


Fig. 4. Response of tobacco leaf tissue to purified $HrpN_{Ech}$. Leaf panel 1 was infiltrated with a suspension of purified $HrpN_{Ech}$ at a concentration of 336 $\mu g/ml$ in 5 mM morpholinoethanesulfonic acid, pH 6.5. Panel 2 was infiltrated with buffer alone. The tissue in panel 1 collapsed 18 hr later. The leaf was photographed, 24 hr after infiltration, with a cross-polarized transilluminator, which enhances black and white visualization by making necrotic, desiccated areas that are typical of the hypersensitive response appear black.

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from kanamycin-resistant colonies and transformed into E. coli DH5a, with selection for kanamycin resistance. Plasmids containing Tn5-gusA1 were analyzed by restriction mapping. Two independent insertions in an 0.82-kb ClaI fragment internal to hrpN_{Ech} were chosen for further study. The precise location and orientation of these insertions was determined by using a primer that hybridizes to Tn5-gusA1 DNA upstream of gusA to sequence into the disrupted E. chrysanthemi DNA (Fig. 1). E. coli DH5α(pCPP2142) cells carrying the Tn5gusA1 insertion at nucleotide 439 of the hrpN_{Ech} ORF (with gusA and hrpN_{Ech} in the same orientation) produced dark blue colonies indicative of \(\beta\)-glucuronidase activity on LM agar (Hanahan 1983) supplemented with 5-bromo-4-chloro-3indolyl \beta-D-glucuronide (data not shown). Whether gusA was expressed from an E. chrysanthemi promoter or the vector lac promoter was not determined. The hrpN_{Ech} 439::Tn5-gusA1 and hrpN_{Ech}546::Tn5-gusA1 mutations were marker-exchanged into the genome of E. chrysanthemi CUCPB5006 (ApelABCE) to produce mutants CUCPB5046 and CUCPB-5045, respectively. Neither of the hrpN_{Eth} mutants elicited a visible reaction in tobacco leaves (Fig. 5).

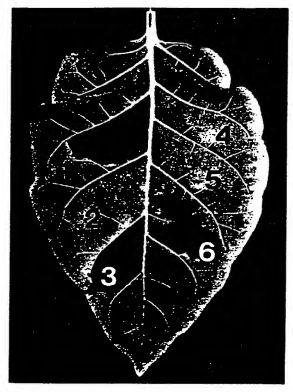


Fig. 5. Tobacco leaf showing that Erwinia chrysanthemi hrpN mutants do not elicit the hypersensitive response unless complemented with $hrpN^*$ pCPP2174. Bacteria were suspended at a concentration of 5×10^3 cells per milliliter in 5 mM morpholinoethanesulfonic acid, pH 6.5, and infiltrated into a tobacco leaf. The leaf was photographed 24 hr later under cross-polarized transillumination. as in Figure 4. 1, E. chrysanthemi CUCPB5006 ($\Delta pelABCE$); 2, CUCPB5045 ($\Delta pelABCE$ hrpN_{Ech}546:: Tn5-gusA1 derivative of CUCPB5006); 3, CUCPB5045(pCPP2174); 4, buffer alone: 5, CUCPB5046 ($\Delta pelABCE$ hrpN_{Ech}439:: Tn5-gusA1 derivative of CUCPB5006); 6, CUCPB5046(pCPP2174).

E. chrysanthemi $hrpN_{Ech}$ mutations can be complemented in trans with $hrpN_{Ech}$ but not with $hrpN_{Ea}$.

The presence of a typical rho-independent terminator just downstream of the hrpN_{Ech} ORF suggested that mutations in the gene would not have polar effects on any other genes and that the HR elicitation phenotype should be restored by an hrpN_{Ech} subclone. Because pCPP2172 carried 2 kb of E. chrvsanthemi DNA in addition to hrpN_{Ech}, we constructed a precise subclone of the gene for this purpose. Oligonucleotides were used to amplify the $hrpN_{Ech}$ ORF by polymerase chain reaction and to introduce terminal NcoI and XhoI sites. The introduction of the restriction sites resulted in changing the second residue in the protein from glutamine to valine and adding a leucine and a glutamic acid residue to the C terminus. The resulting DNA fragment was ligated into XhoI- and NcoI-digested pSE280. creating pCPP2174, in which hrpN_{Ech} was under control of the vector tac promoter. E. chrysanthemi CUCPB5045(pCPP2174) and CUCPB5046(pCPP2174) possessed HR elicitor activity (Fig. 5). HR elicitor activity could also be restored to these mutants by pCPP2142 and pCPP2172, but not by pCPP2141 (data not shown). Thus, the production of HrpN_{Eth} is essential for elicitation of the HR by E. chrysanthemi CUCPB5006.

The feasibility of testing the interchangeability of the hrpN genes of E. chrysanthemi and E. amylovora was supported by the observation that HR elicitation activity could be restored to hrpN mutants in each species (E. chrysanthemi CUCPB-5045 and E. amylovora Ea321T5) by their respective hrpN* subclones (pCPP2142 and pCPP1084). pCPP2142 was used for this purpose because preliminary immunoblot experiments indicated that the level of hrpN_{Ech} expression by this plasmid, though relatively high, most closely approximated the expression of the native hrpN gene in E. amylovora. However, despite good heterologous expression of the hrpN genes, HR elicitation activity was not restored in either E. amylovora Ea321T5(pCPP2142) or E. chrysanthemi(pCPP1084) (data not shown). Thus, the genes do not appear to be functionally interchangeable.

E. chrysanthemi $hrpN_{Ech}$ mutants have a reduced ability to incite lesions in without chicory.

The hrpN_{Ect}439::Tn5-gusA1 mutation was marker-exchanged into the genome of wild-type strain AC4150. The resulting mutant, CUCPB5049, was analyzed for its virulence in witloof chicory. Leaves were inoculated at small wounds with

Table 1. Effects of hrpN_{Eth} mutation on the ability of Erwinia chrysanthemi to incite lesions on witloof chicary leaves

Strain	Number of lesions per 20 inoculations*	Size of lesions (mm², mean ± SD) ^b
AC4150 (wild type) CUCPB5049 (hrpN _{Ech} 429;	16	80 ± 55
TnS-gusAI)	8¢	89 ± 42

^a Each witloof chicory leaf was inoculated at two equivalent sites with 2 × 10⁴ bacterial cells: one site received the hrpN_{Eh} mutant, the other the parental wild-type strain: lesions were indicated by browning and maceration around the site of inoculation.

b Product of the length and width of the lesion.

⁶ Different from the wild-type strain (P < 0.05), as determined by the McNemar test (Conover 1980).

Fig. 6. African violet leaves showing rapid necrosis elicited by HrpN_{Ech} and HrpN²_{Cob} Pel-deficient strains of Erwinia chrysanthemi. Leaves were inoculated with bacteria at a concentration of 3 × 10⁸ cells per milliliter in 5 mM morpholinoethanesulfonic acid, pH 6.5, or purified HrpN_{Ech} at a concentration of 336 μg/ml and photographed 24 hr later under cross-polarized transillumination, as in Figure 4. Buffer controls elicited no visible response (not shown). 1, E. chrysanthemi CUCPB5006 (ΔρεΙΑΒΕΕ): 2, CUCPB5030 (outD::TnphoA derivative of CUCPB-5006); 3, HrpN_{Ech}; 4. (left) CUCPB5045 (ΔρεΙΑΒΕΕ hrpN_{Ech}546::Tn5-gusA1 derivative of CUCPB5063 (ΔρεΙΑΒΕΕ outD::TnphoA hrpN_{Ech}546::Tn5-gusA1 derivative of CUCPB5006).

72 h after inoculation. The mutations did not abolish the pathogenicity of E. chrysanthemi, but they significantly reduced the number of successful lesions (Table 1). However, the $hrpN_{Ech}$ mutation had no significant effect on the size of the lesions produced in successful infections.

Elicitation of a rapid necrosis in several plants by E. chrysanthemi is dependent on $HrpN_{Ech}$.

To determine whether E. chrysanthemi could cause an HrpN_{Ech}-dependent necrosis in plants other than tobacco, a variety of plants were infiltrated with purified HrpN_{Eth} or inoculated with Pel-deficient E. chrysanthemi strains. The strains used were CUCPB5006; its hrpN_{Ech}546::Tn5-gusA1 derivative, CUCPB5045; CUCPB5030 (ApelABCE outD:: TnphoA); and its hrpN_{Ech} 546:: Tn5-gusA1 derivative, CUCPB-5063. The results for African violet are shown in Figure 6, and results for all plants are summarized in Table 2. They yield several general observations. Plants responded either to both isolated HrpN_{Ech} and hrpN_{Ech} bacteria or to neither. Plants that responded to either treatment produced a nonmacerated, HR-like necrosis that developed between 12 and 24 h after infiltration. hrpN_{Ech} mutants failed to elicit a response in any of the plants. The out mutation had no apparent influence on the responses elicited in the plants tested, indicating that residual Pel isozymes or other proteins traveling the Out pathway were not involved in producing the HR-like necrosis. The results argue that $HrpN_{Ech}$ is the only elicitor of the HR produced by E. chrysanthemi.

DISCUSSION

E. chrysanthemi was found to produce a protein with many similarities to the harpin of E. amylovora. The two proteins share significant amino acid sequence identity, similar physical properties, and the ability to elicit the HR in a variety of plants. Mutations in the $hrpN_{Ech}$ gene indicate that, as with E. amylovora, harpin production is required for elicitation of the HR. Furthermore, both harpins contribute to bacterial pathogenicity, albeit to different degrees. $HrpN_{Ea}$ is essential for E. amylovora to produce symptoms in highly susceptible, immature pear fruit (Wei et al. 1992), whereas $HrpN_{Ech}$ merely increases the frequency of successful E. chrysanthemi infections in susceptible witloof chicory leaves. Nevertheless, the finding that harpins play some role in the pathogenicity of

Table 2. Elicitation of necrosis in various plants by HrpN_{Ech} and by Erwinia chrysanthemi strains variously deficient in Pel production and HrpN_{Ech} production

Plant	HrpN _{Ech} ⁴	CUCPB5006 (ΔpelABCE) ^b	CUCPB5045 (\DelABCE hrpN _{Ech} 546:: Tn5-gusA1)	CUCPB5030 (ApelABCE outD::TnphoA)	CUCPB5063 (\DelABCE outD::TnphoA hrpN _{Ech} 546:: TnS-gusA1)
Tobacco	+	+	_	+	-
Tomato	+	+	_	+	_
Pepper	+	+	_	+	_
African violet	+	+	_	<u>.</u>	_
Petunia	+	+	_	<u>.</u>	-
Pelargonium	+	+	_	<u>.</u>	_
Squash		_	-	<u>.</u>	_
Zinnia	_	_	_		_

Leaves on plants were infiltrated with HrpN_{Ech} at a concentration of 336 μg/ml and observed macroscopically 24 hr later for necrosis and collapse of the infiltrated area (+) or absence of any response (-).

b Leaves on plants were infiltrated with bacteria at a concentration of 5 × 108/ml and scored for responses as described above.

such disparate pathogens suggests that these proteins have a conserved and widespread function in bacterial plant pathogenesis. We will consider below $HrpN_{Ech}$ with regard to the protein secretion pathways, extracellular virulence proteins, and wide host range of E. chrysanthemi.

E. chrysanthemi secretes proteins by multiple, independent pathways. Several protease isozymes are secreted by the Secindependent (ABC-transporter, or Type I) pathway; pectic enzymes and cellulase are secreted by the Sec-dependent (general secretion, or Type II) pathway: and. HrpN_{Eth} is likely to be secreted by the Sec-independent Hrp (Type III) pathway (Salmond 1994). The expectation that $HrpN_{Ech}$ is secreted by the Hrp pathway is supported by several lines of indirect evidence: (i) Hrp secretion pathway mutants have revealed that other members of this class of glycine-rich, heat-stable elicitor proteins—the E. amylovora HrpNEa. P. syringae pv. syringae HrpZ, and P. solanacearum PopA1 proteins-are secreted by this pathway (He et al. 1993; Wei and Beer 1993; Arlat et al. 1994); (ii) mutation of the E. chrysanthemi homolog of an E. amylovora gene involved in HrpNE secretion abolishes the ability of E. chrysanthemi to elicit the HR, whereas mutation of the Out (Type II) pathway of E. chrysanthemi does not abolish the HR: and (iii) HrpN_{Ech} appears to be the only HR elicitor produced by E. chrysanthemi (as discussed further below), suggesting that the effect of the putative hrp secretion gene mutation is on $HrpN_{Ech}$. Our attempts to directly demonstrate hrp-dependent secretion of HrpN Ech have been thwarted by the apparent instability of the protein in E. chrvsanthemi. Using the cell fractionation and immunoblotting procedures of He et al. (1993) and polyclonal anti- $HrpN_{Ea}$ antibodies that cross-react with $HrpN_{Ech}$ (Wei et al. 1992), we have observed the presence of HrpN_{Ech} in the cellbound fraction of E. chrysanthemi (D. W. Bauer, unpublished). However, some culture preparations as expectedly lack the protein, and no preparations reveal accumulation of the protein in the culture supernatant fraction. ... :s possible that HrpN_{Eth} aggregates upon secretion and therefore precipitates from the medium. It is interesting that several of the Yersinia spp. Yop virulence proteins aggregate in the medium upon secretion via the Type III pathway (Michiels et al. 1990). Similarly, HrpN_{Ee} has a propensity to form aggregates or to associate with an insoluble membrane fraction (Wei et al. 1992).

It is significant that there is little difference in the plant interaction phenotypes of E. chrysanthemi mutants deficient in either $\operatorname{HrpN}_{Ech}$ or a putative component of the Hrp secretion pathway (Bauer et al. 1994). Both mutations abolish the ability of Pel-deficient strains to elicit the HR , and they both reduce the frequency of successful infections incited by fully pectolytic strains in witloof chicory leaves without affecting

Table 3. Bacterial strains and plasmids used in this study

Designation	Relevant characteristic	Reference or source
Escherichia coli		
ED8767	supE44 supF58 hsdS3(rama) recA56 galk2 galT22 metB1	Sambrook et al. 1989
DH5a	supE44 AlacU169 (\$80 lacZAM15) hsdR17 recA1 endA1 gyrA96 thi-1	Hanahan 1983
	relAI Nal'	Life Technologies, Inc., Grand Island, NY
DH10B	mcrA Δ(mrr-hsdRMS-mcrBC) φ80 lacZΔM15 ΔlacX74 deoR recA1	Grant et al. 1990
	endA1 araD139 Δ(ara, leu)7697 galU galK rpsL nupG	Life Technologies, Inc.
Erwinia chrysanthemi		
EC16	Wild-type strain	Burkholder et al. 1953
AC4150	Spontaneous Nal' derivative of EC16	Chatterjee et al. 1983
CUCPB5006	$\Delta(pelB \ pelC)$:: 28bp $\Delta(pelA \ pelE)$ derivative of AC4150	He and Collmer 1990
CUCPB5030	outD::TnphoA derivative of CUCPB5006	Bauer et al. 1994
CUCPB5045	hrpN _{Ech} 546::TnS-gusA1 derivative of CUCPB5006	This work
CUCPB5046	hrpN _{Ech} 439:: TnS-gusA1 derivative of CUCPB5006	This work
CUCPB5063	hrpN _{Ech} 546::Tn5-gusA1 derivative of CUCPB5030	This work
CUCPB5049	hrpN _{Ech} 439::Tn5-gusA1 derivative of AC4150	This work
Erwinia amylovora		
Ea321	Wild type	ATCC 49947
Ea321T5	hrpN _{Ea} ::TnStac1 derivative of Ea321	Wei et al. 1992
Plasmids and phage		
pBluescript II SK(-)	Amp'	Stratagene, La Jolla, CA
pCPP19	Cosmid vector, Sp'/Sm'	D. W. Bauer
pUC119	Amp' plasmid vector	Vieira and Messing 1987
pSE280	Amp' plasmid vector with superpolylinker downstream of tac promoter	Brosius 1989
pCPP2030	pCPP19 carrying E. chrysanthemi DNA hybridizing with E. amylovora hrp genes in pCPP1033	Bauer et al. 1994
pCPP1084	pBluescript M13+ carrying hrpN _{Ech} on 1.3-kb Hindlll fragment	Wei et al. 1992
pCPP2157	pCPP19 carrying E. chrysanthemi DNA hybridizing with E. amylovora hrpN	This work
pCPP2142	8.3-kb Sall subclone from pCPP2157 in pUC119	This work
pCPP2141	3.1-kb Pst1 subclone from pCPP2157 in pBluescript II SK(-) hrpN _{Ech} in the orientation opposite that of the vector lac promoter	This work
pCPP2172	3.1-kb Pstl subclone from pCPP2157 in pBluescript II SK(-) hrpN _{Ech} in same orientation as vector lac promoter	This work
pCPP2174	1.0-kb hrpN _{Ech} polymerase chain reaction product cloned in Ncol- HindIII sites of pSE280	This work
λ::Tn5-gusA1	Tn5 derivative for generating transcriptional fusions with uidA reporter: Kan', Tet'	Sharma and Signer 1990

Amp' = ampicillin resistance; Kan' = kanamycin resistance; Nal' = nalidixic acid resistance; Sm' = streptomycin resistance; Sp' = spectinomycin resistance; Tet' = tetracycline resistance.

the size of the macerated lesions that do develop. This pattern contrasts with that observed in mutations affecting Pel isozymes and the Out pathway. Virulence, as measured by maceration, is merely reduced by individual pel mutations, whereas it is abolished by out mutations. This is because multiple Pel isozymes (and possibly other enzymes) contribute quartitatively to virulence, but all of the Pel isozymes appear to be dependent on the Out pathway for secretion from the bacterial cell. The simplest interpretation of the observations with E. chrysanthemi hrp mutants is that HrpN_{Ech} is the only protein traveling the Hrp pathway that has a detectable effect on the interaction of E. chrysanthemi EC16 with the plants tested.

The primacy of $HrpN_{Ech}$ in the E. chrysanthemi Hrp system is further supported by the observations that $hrpN_{Ech}$ mutants failed to elicit necrosis in any of the several plants tested and that all plants responding with apparent hypersensitivity to HrpN+ch strains also responded to isolated HrpN-ch. Several of the plants sensitive to HrpN Ech are also susceptible to bacterial soft rots. This is particularly significant for African violet, whose interactions with E. chrysanthemi have been extensively studied (Barras et al. 1994). Thus, $HrpN_{Ech}$ elicits HRlike responses in plants that are susceptible to E. chrysanthemi infections under appropriate environmental conditions. The significance of this for the wide host range of the bacterium requires further investigation, and virulence tests with hrpN_{Ech} mutants and additional susceptible plants are needed to determine the general importance of HrpNEch and the Hrp system in E. chrysanthemi. For example, our present data do not address the possibility that other proteins secreted by the Hrp pathway, which are not elicitors of the HR in the plants we tested, may contribute to pathogenesis in hosts other than witloof chicory.

An important question is whether bacteria expressing heterologous harpins will be altered in pathogenicity. The hrpN genes of E. chrysanthemi and E. amylovora are particularly attractive for experiments addressing this because of the similarity of the harpins and the dissimilarity of the diseases produced by these bacteria. Unfortunately, attempts to restore the HR phenotype to E. chrysanthemi and E. amylovora hrpN mutants with heterologous hrpN+ subclones failed. Since the hrpN genes in each subclone successfully complemented hrpN mutations in homologous bacteria and were expressed in heterologous bacteria, the problem is most likely the secretion of the harpins by heterologous Hrp systems. A similar problem has been encountered with heterologous secretion of Pel and cellulase via the Out pathway in E. chrysanthemi and E. carotovora, species that are more closely related to each other in this rather heterogeneous genus than E. chrysanthemi and E. amylovora are (He et al. 1991; Py et al. 1991).

In conclusion, two classes of proteins contribute to the pathogenicity of *E. chrysanthemi*—a single harpin and a battery of plant cell wall-degrading pectic enzymes. The observation that such a highly pectolytic organism also produces a harpin suggests the fundamental importance of harpins in the pathogenicity of gram-negative bacteria. The observation that an *hrpN_{Ech}*::Tn5-gusAI mutation reduced the ability of a fully pectolytic strain of *E. chrysanthemi* to initiate lesions in susceptible chicory leaves, but did not reduce the size of lesions that did develop, suggests that HrpN_{Ech} contributes specifically to an early stage of pathogenesis. An attractive pos-

sibility is that $HrpN_{Ech}$ releases nutrients to the apoplast for bacterial nutrition before the *pel* genes are fully expressed (Collmer and Bauer 1994). Patterns of *pel* and $hrpN_{Ech}$ expression in planta will likely yield further clues to the role of the *E. chrysanthemi* harpin in soft-rot pathogenesis.

MATERIALS AND METHODS

Bacterial strains, plasmids, and culture conditions.

Bacterial stains and plasmids are listed in Table 3. E. chrysanthemi was routinely grown in King's medium B (King et al. 1954) at 30° C. E. coli in LM medium (Hanahan 1983) at 37° C, and E. amylovora in Luria-Bertani medium at 28-30° C. The following antibiotics were used in selective media in the amounts indicated (in µg/ml), except where noted: ampicillin (100), kanamycin (50), spectinomycin (50), and streptomycin (25).

General DNA manipulations.

Plasmid DNA manipulations, colony blotting, and Southern blot analyses were performed by standard techniques (Sambrook et al. 1989). Deletions for sequencing were constructed with the Erase-a-Base kit (Promega, Madison, WI). Doublestranded DNA sequencing templates were prepared with Qiagen Plasmid Mini Kits (Chatsworth, CA). Sequencing was performed with the Sequenase Version 2 kit (U.S. Biochemical, Cleveland, OH). The Tn5-gusA1 insertion points were determined on an Automated DNA Sequencer (model 373A. Applied Biosystems, Foster City, CA) by the Cornell Biotechnology Center. DNA sequences were analyzed with the Genetics Computer Group Sequence Analysis Software Package (Devereaux et al. 1984). Comparison of HrpN_{Ech} and HrpN_{Ea} by the Gap program was done with a gap weight of 5.0 and a gap length weight of 0.3. Marker exchange mutagenesis was performed as previously described (Bauer et al. 1994). The oligonucleotide used to determine the location of Tn5-gusA1 insertions in hrpN_{Ech} was TGACCTGCAGCC-AAGCTTTCC. The oligonucleotide used as the first primer to amplify the hrpN_{Ech} ORF and introduce an NcoI site at the 5' end of the gene was AGTACCATGGTTATTACGATCAAA-GCGCAC; the one used as the second primer to introduce an XhoI site at the 3' end of the gene was AGATCTCGAGGG-CGTTGGCCAGCTTACC. Primers were synthesized by Integrated DNA Technologies (Coralville, IA).

Protein manipulations.

HrpN_{Ech} was purified from E. coli DH5α(pCPP2172) cultures grown at 30° C to stationary phase in 50 ml of Terrific Broth (Sambrook et al. 1989) supplemented with ampicillin at a concentration of 200 µg/ml. Cells were lysed by lysozyme treatment and sonication as previously described (Sambrook et al. 1989). The lysate pellet was washed twice with 9 vol of lysis buffer containing 0.5% Triton X-100 and 10 mM EDTA, pH 8.0, and 0.1 mM phenylmethylsulfonyl fluoride (PMSF); the lysate was reharvested each time by centrifugation at $12,000 \times g$ for 15 min. The pellet was resuspended in 2.0 ml of lysis buffer containing 0.1 mM PMSF, dissolved by the addition of 2.5 ml of 8 M guanidine-HCl in lysis buffer, and then diluted with 5.0 ml of water. The protein solution was dialyzed in SpectraPor #1 dialysis tubing against 2.0 liters of 5 mM morpholinoethanesulfonic acid (MES), pH 6.5. containing 0.05 mM PMSF. The precipitate that formed during dialysis and the solution were centrifuged for 15 min at $4,300 \times g$. The pellet was washed once with 10 ml of a solution containing 5 mM MES, pH 6.5, and 0.1 mM PMSF and then resuspended in 2.0 ml of the same buffer. Protein concentrations of homogeneous suspensions were determined following dissolution in the reagents of the dye-binding assay of Bradford (1976). Proteins in crude cell lysates or following purification were resolved by electrophoresis through an SDS 12% polyacrylamide gel and visualized by staining with Coomassie Brilliant Blue R. The N-terminal sequence of purified HrpN_{Ech} was determined at the Cornell University Biotechnology Program Protein Analysis Facility.

Plant assays.

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> For HR assays, tobacco (Nicotiana tabacum L. cv. Xanthi). tomato (Lycopersicon esculentum Mill. cv. Sweet 199), pepper (Capsicum annuum L. cv. Sweet Hungarian), African violet (Saintpaulia ionantha H. Wendl. cv. Paris), petunia (Petunia grandiflora Juss. cv. Blue Frost), pelargonium (Pelargonium hortorum Bailey), winter squash (Cucurbita maxima Duchesne), and zinnia (Zinnia elegans Jacq.) plants were grown under greenhouse conditions or purchased at a local garden shop and then maintained in the laboratory at room temperature, with incident daylight supplemented with a 500-W halogen lamp. Witloof chicory (Cichorium inrybus L.) was purchased as "Belgian endive" heads from a local supermarket. Bacterial inoculum was prepared and delivered as previously described (Bauer et al. 1994). Briefly, to assay soft-rot pathogenesis, 5 µl of inoculum was applied to a small wound in detached chicory leaves; to assay for HR elicitation, inoculum was infiltrated with a needle-less plastic syringe into leaves on plants.

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The RsmA⁻ Mutants of *Erwinia carotovora* subsp. *carotovora* Strain Ecc71 Overexpress *hrpN*_{Ecc} and Elicit a Hypersensitive Reaction-like Response in Tobacco Leaves

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Erwinia carotovora subsp. carotovora wild-type strain Ecc71 does not elicit the hypersensitive reaction (HR) in tobacco leaves. By mini-Tn5-Km and chemical mutagenesis we have isolated RsmA mutants of Ecc71 that produce high basal levels of pectate lyases, polygalacturonase, cellulase, and protease; they also are hypervirulent. The RsmA mutants, but not their parent strains, elicit an HRlike response in tobacco leaves. This reaction is characterized by the rapid appearance of water soaking followed by tissue collapse and necrosis. The affected areas remain limited to the region infiltrated with bacterial cells, and the symptoms closely resemble a typical HR, e.g., the reactions caused by Pseudomonas syringae pv. pisi. Moreover, low concentrations of cells of the mini-Tn5-Km insertion RsmA mutant, AC5070, infiltrated into tobacco leaf tissue prevent elicitation of the rapid necrosis by AC5070 or by P. syringae pv. pist. Elicitation of the HR-like response by the mutants is not affected by the deficiency of N-(3oxohexanoyl)-L-homoserine lactone, the cell density (quorum) sensing signal. Cloning and sequence analysis have disclosed that E. carotovora subsp. carotovora strain Ecc71 possesses a homolog of B. chrysanthemi hrpN known to encode an elicitor of the HR; the corresponding Ecc71 gene is designated hrpN_{Ecc} Northern (RNA) blot data show that the level of hrpNE mRNA is considerably higher in the RsmA* mutants than in the RsmA* strains. Moreover, a low copy plasmid carrying the rsmA* allele severely reduces the level of the hrpN see transcripts in the RsmA" mutants. These constructs, like the RsmA* E. carotovora subsp. carotovora strains, do not elicit the HRlike response. These data taken along with the effects of rsmA on exoenzyme production and pathogenicity (A. Chatterjee et al., 1995, Appl. Environ. Microbiol. 61:1959-1967) demonstrate that this global regulator gene plays a critical role in plant interaction of E. carotovora subsp. carotovora.

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Nucleotide and/or amino acid sequence data is to be found at GenBank as accession number L78834.

Additional keywords: derepressed mutant, incompatible interactions, soft-rotting bacteria.

Many gram-negative phytopathogenic bacteria, when infiltrated into a nonhost plant such as tobacco, cause localized necrosis, generally known as the hypersensitive reaction (HR) (Goodman and Novacky 1994). A typical HR is characterized by the rapid collapse of the leaf tissue followed by necrosis of the collapsed area. Erwinia carotovora subsp. carotovora and many other soft-rotting bacteria are unusual in that they do not elicit a typical HR when infiltrated into tobacco leaves. The inability of these bacteria to elicit the HR has been attributed to the production of pectolytic enzymes that are presumed to suppress the HR. The recent finding of Collmer and his associates that a mutant strain of E. chrysanthemi deficient in the synthesis of the major pectate lyase (Pel) isozymes, but not the pectolytic parent, can elicit the HR (Bauer et al. 1994) is certainly consistent with this hypothesis. In fact, both genetic and biochemical data (Bauer et al. 1995) demonstrate that E. chrysanthemi, like many other gram negative bacteria, possesses hrp genes including hrpN, which encodes an elicitor of the HR. These data and the results of Southern blot hybridizations of Laby and Beer (1992) support the idea that softrotting Erwinia possess hrp genes, but a sustained expression of hrp genes of these Erwinia species in incompatible hosts may not occur at a level required for elicitation of the

We have initiated studies to clarify the genetic regulation of the production of the HR and disease symptoms by E. carotovora subsp. carotovora. We previously reported that a mini-Tn5-Km insertion RsmA mutant of E carotovora subsp. carotovora is derepressed in extracellular enzyme production and it is hypervirulent (Chatterjee et al. 1995; Cui et al. 1995). A mutant of similar phenotype was also generated by chemical mutagenesis. The data presented here show that these mutants elicit responses in tobacco leaves that are similar to those in a typical HR and that they do not require the cell density sensing signal, N-(3-oxohexanoyl)-L-homoserine lactone (OHL) to cause this reaction. Additionally, our findings disclose the presence of a homolog of the hrpN_{E0} gene in E. carotovora subsp. carotovora strain Ecc71 and show that expression of this gene is negatively controlled by rsmA.

RESULTS

RsmA mutants of E. carotovora subsp. carotovora elicit responses in tobacco leaves that resemble the HR.

Previously (Chanerjee et al. 1995; Cui et al. 1995), we have described the isolation procedure as well as some of the characteristics of E. carotovora subsp. carotovora strain AC5070, the mini-Tn5-Km insertion RsmA- mutant (rsm = regulator of secondary metabolites). Since AC5070 overproduces peciate lyases, polygalacturonases, protease, and cellulase, and is hypervirulent, it was of interest to examine the responses it could elicit in tobacco leaves, wherein wild-type E. carosovora subsp. carotovora does not cause tissue necrosis in 24 to 48 hr. As shown in Figure 1, cells of AC5070 infiltrated into tobacco leaves produced symptoms similar to those caused by P. syringae pv. pisi, known to elicit the HR. The lowest concentration of AC5070 that elicited an HR-like response was approximately 2 × 10⁸ cells/ml. The visible symptoms, i.e., water soaking followed by tissue collapse, appeared within 24 h after the infiltration. By 24 h the inoculation sites developed necrosis, culminating in tissue desiccation. These responses, as in the typical HR, invariably remained confined to the area infiltrated with bacterial cells. Infiltration with cells of RsmA* E. carotovora subsp. carotovora grown in Luria-Bertani (LB) agar did not produce visible lesions; however, after 5 to 6 days the infiltrated sites became chlorotic.

By ethyl methane sulfonate (EMS) mutagenesis of *E. carotovora* subsp. *carotovora* strain AC5006, we isolated a mutant, AC5041, that, like AC5070, overproduces pectate lyases, polygalacturonases, protease, and cellulase (Fig. 2). In addi-

Fig. 1. Symptoms produced in tobacco leaves by Erwinia carotovora subsp. carotovora AC5047 and its RsmA" mutant, AC5070. Cell suspensions containing about 2 × 10^t CFU/ml were infiltrated into each leaf segment. A, AC5047; B, AC5070; C, Pseudomonas syringue pv. pisi Psp1; and D, water. Picture was taken 24 h after infiltration.

tion, the mutant is hypervirulent in that it caused more severe maceration in celery petioles than the parent RsmA⁺ strain (Fig. 3). The derepressed mutant, AC5041, but not its parent strain, induced the HR-like response in tobacco leaves (data not shown).

Prevention of the HR-like response.

It has been reported that P. syringae pv. pisi prevents the HR when it is preinoculated in tobacco leaves at a lower concentration (5×10^5) and later challenged with an HR-inducing concentration (5×10^6) at the same site (Novacky et al. 1973). Similarly, we have noticed that preinfiltration of tobacco leaves with AC5070 (10^5 CFU/ml) prevented the appearance of water soaking and necrosis upon reinoculation at the same

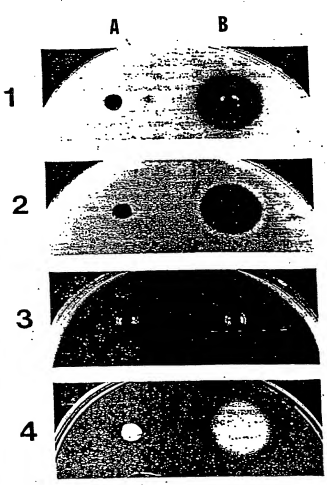


Fig. 2. Agarose plate assays for 1, pectate lyase (Pel); 2, polygalacturonase (Peh); 3, protease (Pri); and 4, cellulase (Cel) activities of Erwinia carotovora subsp. carotovora AC5006 (A) and its RsmA* munnt AC5041 (B). Bacteria were grown in salts—yeast extract—glycerol medium to saturation. Culture supernatants were diluted twofold in 10 mM Tris-HCI (pH 7.0) buffer and 5 µl of the diluted samples were used for the Pel, Peh, and Cel assays. Thirty microliters of undiluted samples were used for the Pri assay.

site with AC5070 or P. syringae pv. pisi (Fig. 4). After the preinoculation, about 2×10^8 cells of AC5070 were introduced at different intervals. The ability of preinoculated cells to inhibit the HR-like response was apparent by 12 h after inoculation (data not shown), and by 24 h production of the response was completely suppressed.

A B

Fig. 3. Maceration of celery periodes induced by Erwinia carotovora subsp. carotovora AC5006 (A) and its RsmA* mutant AC5041 (B). About 2 × 10⁸ bacterial cells suspended in water were injected into each inoculation site. Inoculated periodes were covered with perroleum jelly and incubated in a moist chamber at 25°C for 24 b.

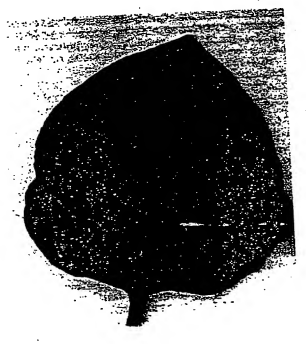


Fig. 4. Prevention of the hypersensitive response symptoms in tobacco leaf by the RimA⁻ mutant of Erwinia carotovora subsp. carotovora. ACS070. Leaf segments were infiltrated with A, water at 0 h; B, Pstadomonas syringae pv. pist Pspl (5 × 10^a CFU/ml) at 24 h; C, ACS070 (2 × 10^a CFU/ml) at 0 h; E, ACS070 (2 × 10^a CFU/ml) at 0 h; E, ACS070 (10⁵ CFU/ml) at 0 h and challenged with Pspl (5 × 10^a CFU/ml) after 24 h; G, ACS070 (10⁵ CFU/ml) at 0 h and challenged with ACS070 (2 × 10^a CFU/ml) after 24 h; and H, Pspl and challenged with ACS070 (2 × 10^a CFU/ml) after 24 h; and H, Pspl (5 × 10^a CFU/ml) at 0 hour. Leaf was photographed 48 h after infiltration.

RsmA⁻ mutants of *E. carotovora* subsp. carotovora elicit the HR-like response in the absence of the cell density sensing signal, OHL.

OHL and its structural analogs are required for the expression of many phenotypes in different bacteria (Fuqua et al. 1994; Salmond et al. 1995; Swift et al. 1994). In E. carotovora subsp. carotovora, OHL controls extracellular enzyme production, pathogenicity, and production of the antibacterial antibiotic, carbapenem (Bainton et al., 1992; Chatterjee et al. 1995; Jones et al. 1993; Pirhonen et al. 1993). We had previously demonstrated that excenzyme overproduction and pathogenicity occurred in the absence of OHL in the RsmAmutant, AC5070 (Chatterjee et al. 1995). To find out if the mutants could elicit the HR-like response in the absence of this cell density sensing signal, we examined the responses induced by OHL deficient derivatives of the RsmA strains. We made the EMS-induced RsmA- mutant OHL deficient by replacing ohll" (previously designated as hell") allele required for OHL biosynthesis, with ohll-Mudl by marker exchange, as we had done with AC5070 (Chatterjee et al. 1995). AC5090 and AC5093, the derivatives of AC5070 and AC5041, respectively, do not produce OHL, as indicated by the Lux bioassay (Chatterjee et al. 1995; data not shown). Figure 5 shows that AC5090 and AC5093 elicited reactions in tobacco leaves that were very similar to those produced by the parent strains as well as by P. syringae pv. pisi.

The RsmA⁻ mutants overexpress $hrpN_{Rec}$ a locus presumed to specify an elicitor of the HR.

Recent studies by S. V. Beer, A. Colliner, and their associates demonstrated that hrpN genes of E amylovora and E. chrysanthemi encode elicitors of the HR and raised the possi-



Fig. 5. Elicitation of the hypersensitive-like response in tobacco leaves by RsmA⁻ mutants of Erwinia carotovora subsp. carotovora and their Ohll' derivatives. Leaf segments were infiltrated with 2 × 10^s CFU/ml of bacterial cells. A, water. B, AC5093, (RsmA⁻, Ohl'); C, AC5090 (RsmA⁻, Ohl'); D, Pseudomonas syringae pv. piri Pspl; E, AC5041 (RsmA⁻, Ohl'); and F, AC5070 (RsmA⁻, Ohl'). Picture was taken 24 h after infiltration.

HIPN _{Ecc} HIPN _{Ech} HIPN _{E4}	MINSIGG-GASIQITIKA-GGNGGHFFSQSSQSSVDKLS	45 43 50
HrpN _{Zec} HrpN _{Zeb} HrpN _{Ze}	NIAEQLSDIMTTMMFMGSMMGGGMSGGLGGLGSSLGGLGGGL STIDKLTSALTSMMFGGALAQGLGASSKGLG DTVNQLAGLLTGMMMMMSMMGGGGLMGGGLGGGLGNGLGGGGGGLGEGLSN 1	87 74 LOO
HrpN _{rec} HrpN _{reb} HrpN _{ra}	-LGGGLGGGLGSSLGSGLGSALGGGLGGALGAA	120 104 149
HIPN _{ECE} HIPN _{ECE} HIPN _{ECE}		167 150 199
HIDN rec HIDN rec HIDN rec	QGVNDNLSAILGNGLSQTKG	203 186 249
HrpN _{Ecc} HrpN _{Ech} HrpN _{Ea}	GAGAFNOLGSTLGMSVGQKAGLQELNNISTHNDSPTRYFVDKEDRGMAKE GAGAFNOLGNAIGMGVGQNAALSALSNVSTHVDGNNRHFVDKEDRGMAKE GPVDYQQLGNAVGTGIGMKAGIQALNDIGTHRHSSTRSFVNKGDRAMAKE	253 236 299
HrpN _{sec} HrpN _{sec}	IGQFMDQYPEVFGKAEYQKDNWQTAKQEDKSWAKALSKPDDDGMTKGSMD IGQFMDQYPEIFGKPEYQKDGWSSPKTDDKSWAKALSKPDDDGMTGASMD IGQFMDQYPEVFGKPQYQKGPGQEVKTDDKSWAKALSKPDDDGMTPASME	303 286 3 4 9
HrpN _{see} HrpN _{seh} HrpN _{se}	KFMKAVGMIKSAIRGDTGNTNLSARGNGGASLGIDAAMIGDRIVNMGLKK KFRQAMGMIKSAVAGDTGNTNLNLRGAGGASLGIDAAVVGDKIANMSLGK QFNKAKGMIKRPMAGDTGNGNLQHAVPVVLRW	353 336 381
HIDN _{EC} HIDN _{EC} HIDN _E	LSS- 356 LANA 340 VIMP 385	th those of E

Fig. 6. Alignment of deduced amino acid sequence of hrpN_{Eoc} of Erwinia carotovora subsp. carotovora strain Ecc71 (HrpN_{Eoc}) with those of E chrysanthemi EC16 (HrpN_{Eoc}) and E. amylovora E3321 (HrpN_{Eoc}). Asterisks indicate identical amino acids; single dots indicate conservative substitutions. Numbers at R right indicate amino acid positions in each protein.

bility that hrp genes including hrpN may also occur in other Erwinia species (Bauer et al. 1994; Bauer et al. 1995; Laby and Beer 1992; Wei et al. 1992). Indeed, Southern blot hybridization under moderate stringency conditions with hrpN DNA of E. chrysanthemi (EC16) (Bauer et al. 1995) as the probe disclosed the presence of hrpN sequences in E. carotovora subsp. carotovora strain Ecc71 (data not shown). Subsequently, by screening a library of Ecc71 with the hrpN DNA of E. chrysanthemi, several clones possessing homologous DNA were identified; the corresponding Ecc71 sequences are tentatively designated as hrpNzc. Sequence analysis of the DNA segment that specifically hybridized with the hrpN DNA of E chrysanthemi revealed an 1,068-bp open reading frame whose predicted product has 72.1% similarity and 53.4% identity with the deduced product of hrpN of E. chrysanthemi, and 66.6% similarity and 50.8% identity with the predicted product of hrpN of E. amylovora (Fig. 6).

Northern (RNA) blot analysis was performed with total RNA preparations from the wild-type strain Ecc71, the RsmAmutants, AC5041 and AC5070, and their RsmA+ parents to ascertain if hrpN_{Ecc} expression is derepressed in the RsmA⁻ strains. Bacteria were grown in SYG medium at 28°C to a Klett value of approximately 200 and used for total RNA isolation. A 700-bp Acci-Smal internal fragment of the hrpN_{Eee} was used as the probe. The data (Fig. 7) revealed the presence of 1100-base transcripts in AC5070 and AC5041. By contrast, these transcripts were not detected with RsmA+ strains 71, AC5006 and AC5047. We should note that somewhat higher levels of hrpN_{Ecc} transcripts were present in the mini-Tn.5-Km insertion mutant (AC5070) than in the EMS-induced mutant (AC5041). We do not yet know the reason for this difference. It is possible that AC5041 produces a defective RsmA with a leaky activity, whereas the mini-Tn5-Km insertion mutant does not produce a functional RsmA. It is, however, clear that hrpN_{Eee} transcripts are substantially higher in AC5041 than in its RsmA+ parent, AC5006.

The $rsmA^+$ allele suppresses elicitation of the HR-like response and expression of $hrpN_{\rm geo}$

We have previously described the cloning and characterization of the rsmA gene of E. carotovora subsp. carotovora strain Ecc71 (Chatterjee et al. 1995; Cui et al. 1995). A lowcopy plasmid carrying this gene causes a severe attenuation of pathogenicity and suppresses extracellular enzyme production in E carotovora subsp. carotovora and E. c. subsp. atroseptica; represses pathogenicity, exopolysaccharide production, flagellum production and motility, protease production, and elicitation of the HR by E amylovora; and suppresses extracellular enzyme and antibiotic production by E carotovora subsp. betavascularum (Chatterjee et al. 1995; Cui et al. 1995; Mukherjee et al. 1996a, 1996b). In light of the large array of effects on phenotypes by rsmA, including induction of the HR by E. amylovora, it was deemed worthwhile to examine the effects of the rsmA+ DNA on elicitation of the HR-like response by the mutants. The plasmids pCL1920 and pAKC880 were transformed into AC5041 and AC5070 and the constructs were tested for induction of the HR-like response. Figure 8 shows that AC5041 and AC5070 carrying the cloning vector, pCL1920, elicited reactions in tobacco leaves similar to those caused by P. syringae pv. pisi. By contrast, there was no visible reaction in the leaf segment infiltrated with AC5041

or AC5070 carrying the RsmA⁺ plasmid, pAKC880. These results indicate that multiple copies of rsmA suppress elicitation of the HR-like response in tobacco leaves by AC5041 and AC5070.

Northern analysis was conducted to determine the effect of RsmA plasmid on $hrpN_{Ecc}$ transcription. The data (Fig. 9) show that high levels of $hrpN_{Ecc}$ transcripts were present in cells of AC5041 and AC5070 containing the cloning vector, pCL1920, but the transcripts were not detected in cells carrying the rsmA plasmid, pAKC880.

DISCUSSION

We previously reported that extracellular enzyme production as well as virulence are negatively regulated by rsmA in E. carotovora subsp. carotovora (Chatterjee et al. 1995; Cui et al. 1995; Mukherjee et al. 1996a, 1996b). For example, the inactivation of rsmA by a transposon resulted in overproduction of extracellular enzymes and hypervirulence. Moreover, unlike its parent, the RsmA mutant did not require the cell density sensing signal, OHL, for pathogenesis or extracellular enzyme production. In this report, we have shown that this RsmA mutant and an EMS-induced mutant of a similar phenotype elicited the HR-like response in tobacco leaves, and that the elicitation of this reaction was also not dependent upon OHL. Although we do not yet have direct evidence that the mutations in AC5041 and AC5070 are in the same gene, these strains possess similar phenotypes; e.g., they overproduce extracellular enzymes, they are hypervirulent, and OHL deficiency does not affect the expression of these traits. Moreover, the plasmid carrying rsmA+ DNA suppresses extracellular enzyme production, pathogenicity, and the elicitation

1 2 3 4 5

1100-



Fig. 7. Northern (RNA) blot analysis of httpN_{Ecc} mRNA of Erwinia caratovora subsp. caratovora strains. Each lane contained 20 µg of total RNA. Position of 1100-base transcript is indicated. Lane 1, Ecc71 (wild-type parent, RsmA*); lane 2, AC5006 (RsmA**); lane 3, AC5041 (RsmA**); lane 4, AC5047 (RsmA**); lane 5. AC5070 (RsmA**).

of the HR-like response by the mutants. Also, both the mutants express $hrpN_{Ecc}$ constitutively, although the transcript level is somewhat higher in AC5070 than in AC5041. As these mutants have similar phenotypes, we tentatively classified them as RsmA.

The following lines of evidence strongly suggest that the mutants elicited a typical HR (Goodman and Novacky 1994): (i) the reaction was characterized by a rapid physiological activity (i.e., water movement or water soaking), tissue collapse followed by cell death (necrosis); (ii) the affected areas were limited to the region infiltrated with bacterial cells; (iii) these symptoms were indistinguishable from the symptoms developed by P. syringae pv. pisi, a bacterium known to elicit the typical HR in tobacco leaves; (iv) the response elicited by AC5070 was preventable upon previous infiltration of a low concentration of AC5070 cells and, similarly, prior inoculations with AC5070 cells prevented elicitation of the HR by P. syringae pv. pisi; and (v) while ACS070 and ACS041, their parent strains, and the wild-type strain possess hrpN_{Eee} sequences (data not shown), the expression of hrpN_{Ecc} is derepressed only in the mutants, presumably leading to the production of high levels of a putative elicitor of the HR (see

below).
Our observations support the idea that AC5070 and AC5041 produce an elicitor that triggers the HR-like response

in tobacco leaves. We attribute the manifestation of this response with the mutants, but not with the parents, to the ability of the former to produce high constitutive levels of HrpN_{Eer} an exoenzyme, or both. With regard to the possible role of exoenzymes, it is perhaps significant that pectinases are known to generate elicitors of plant defense responses (Davis et al. 1984; Davis and Ausubel 1989; Keen 1992). Furthermore, Palva et al. (1993) have documented the activation of chitinases and glucanases in tobacco by exoenzymeproducing strains of E. carotovora subsp. carotovora but not by mutants deficient in exoenzyme production. Therefore, one could argue that pectinase overproduction by the RsmA-mutants may induce defense reactions that could culminate in an HR-like response. The inability of the wild-type RsmA* E. carotovora subsp. carotovora strain Ecc71 to elicit this response could be attributed to the lack of extracellular enzyme production in a nonhost tissue, i.e., in a tobacco leaf. However, the hypothesis implicating pectolytic enzymes as elicitors of the HR is difficult to reconcile with the finding of Bauer et al. (1994) that only those mutants of E. chrysantheni that are deficient in major pectate lyases can elicit the HR.

In light of that finding and for the following reasons, we favor the hypothesis that induction of the HR-like response by the mutants may be due to the derepression of a gene encoding an elicitor, such as HrpN_{Ech} or HrpN_{Ech} Collmer and asso-

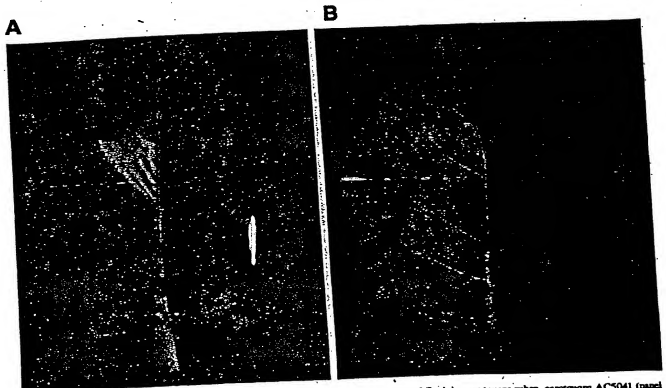


Fig. 8. Elicitation of an hypersensitive-like response in tobacco leaves by the RsmA⁻ mutants of Erwinia carotovora subsp. carotovora AC5041 (panel A) and AC5070 (panel B) carrying the rsmA⁺ plasmid, pAKC380, or the cloning vector, pCL1920. Bacterial suspensions containing about 2 × 10⁸ CFU/ml were infiltrated into each leaf segment. Panel A: A, Pseudomonas syringas pv. pist Psp1; B, AC5041 carrying pAKC380; C, water; D, AC5041 carrying pCL1920. Picture was taken 24 h after infiltration. carrying pCL1920. panel B: A, Psp1; B, AC5070 carrying pAKC380; C, water; D, AC5070 carrying pCL1920.

ciates (Bauer et al. 1994; Bauer et al. 1995) have discovered a gene specifying an elicitor of the HR in the soft-rotting bacterium E. chrysanthemi. The deduced sequence of HrpN Ecr presented here document the occurrence of a homolog of E. chrysanthemi hrpN in E. carotovora subsp. carotovora strain Ecc71. We have found that the mini-Tn5-Km induced RsmAmutant as well as the EMS-induced derepressed mutant possess a substantial level of an approximately 1100-base transcript that specifically hybridizes with the hrpN_{Esc} DNA. By contrast, this transcript is barely detectable in the RsmA+ strains. Moreover, the introduction of the rsmA+ allele into the mutants severely reduces the levels of this transcript and concomitantly abolishes the ability to elicit the HR-like response. These observations indicate that transcription of hrpN_{Exx} is derepressed in the mutants, and that this derepression is due to the inactivation of rsmA. At the moment, since the genes for pectolytic enzymes and hrpNex are both derepressed in the RsmA mutants, we have to entertain the possibility that the pectolytic enzymes could also contribute to the hypersensitive necrosis of tobacco leaf tissue. Genetic and biochemical studies have been initiated to determine if hrpN_{Eec} and its putative product are solely responsible for the elicitation of the HR and to clarify the ramifications of hpN_{Scc} regulation in compatible and incompatible interactions of E. carotovora subsp. carotovora.

Fig. 9. Northern (RNA) blot analysis of hrpN_{Eoc} mRNA of Erwinia carotovora subsp. carotovora RsmA* mutants AC5041 and AC5070 carrying the rsmA* plasmid, pAKC880, or the cloning vector. pCL1920. Each lane contained 20 pg of total RNA. The position of 1100-base tran-Each lane contained 20 pg of total KIVA. The position of 1100-base transcript is indicated. Lane 1, AC\$070 carrying pCL1920; lane 2, AC\$070 carrying pAKC880; lane 3, AC\$041 carrying pCL1920; lane 4, AC\$041 carrying pAKC880.

MATERIALS AND METHODS

Bacterial strains and media.

Bacterial strains and plasmids are described in Table 1. E. carotovora subsp. carotovora strains were routinely grown in LB and P. syringae pv. pisi on King's B (King et al. 1954) agar media at 28°C. Minimal salts plus sucrose (0.2%) agar, nutrient gelatin (NG) agar, polygalacturonate-yeast extract agar (PYA) and salts-yeast extract-glycerol (SYG) media have been described previously (Barras et al. 1987; Chatterjee 1980; Murata et al. 1991). When required, antibiotics were added at the indicated concentrations in micrograms per milliliter: spectinomycin (Spc), 50; tetracycline (Ic), 10; Ampicillin (Ap), 50 and Kanamycin (Km), 50. The composition of agarose media for semiquantitative assays of enzymatic activities has been described in Chatterjee et al. (1995).

Enzyme assays.

The preparation of enzyme samples for assays as well as the assay procedures were described previously (Murata et al. 1991; Chanterjee et al. 1995). The volumes of enzyme samples used in the assays are indicated in the figure legends.

Bioluminescence assay for OHL.

The procedure described by Chanterjee et al. (1995) was followed.

Recombinant DNA techniques.

Standard procedures were followed in DNA isolation, transformation and electroporation of bacteria, restriction digests, gel electrophoresis, DNA ligation, and Southern blot analysis (Sambrook et al. 1989). Restriction and modifying enzymes were obtained from Promega Biotech (Madison, WI).

Isolation of RsmA-mutants.

The procedure used for the isolation of AC5070 by mini-Tn5-Km has been described (Chatterjee et al. 1995). AC5041 was isolated by EMS mutagenesis of AC5006. Mutagenesis was carried out according to the protocol of Miller (1972). The bacterial cells were incubated with EMS for a period that yielded less than 5% survival. The putative RsmA- mutants were identified by their ability to overproduce protease, cellulase, and pectolytic enzymes in agar plate assays (Chatterjee et al. 1995).

Inactivation of the ohl locus by MudI mutagenesis.

The plasmid, pAKC852, carrying the 9.7-kh ohl+ DNA of E. carotovora subsp. carotovora strain Ecc71 was mutagenized with MudI1734 following the procedure of Castilho et al. (1984). Briefly, pAKC852 was transformed into the lysogenic Escherichia coli strain POI1734. The strain carrying the Ohl* plasmid was heat-induced to lyse. The lysate was used to transduce E. coli M8820, and the Tc'Km' transductants were screened for OHL production by means of the plate assay procedure described in Chatterjee et al. (1995). Plasmids were isolated from M8820 colonies that could no longer activate the lux operons in pHV200L

Construction of bacterial strains by marker exchange.

The construction of AC5090, the Ohl derivative of AC5070, has been described (Chatterjee et al. 1995). To isolate AC5093, the Ohl mutant of AC5041, the plasmid (pAKC863) carrying inactivated ohll-Mudl was transferred into AC5041 by means of the helper plasmid, pRK2013. Transconjugants were selected on minimal salts plus sucrose agar supplemented with Km. Colonies that were KmTc were tested for the Ohl phenotype. AC5093 was selected for further studies.

Plant tissue maceration.

The celery periole assay was previously described (Murata et al. 1991). The extent of tissue maceration was estimated visually.

Infiltration of tobacco leaves.

Erwinia species were grown on LB agar and P. syringae pv. pisi was grown on King's B agar overnight at 28°C and cells were resuspended in water. Strains carrying plasmids were grown on LB agar containing spectinomycin and cells suspended in a 50 µg/ml spectinomycin solution in water. Young, fully expanded third and fourth leaves of about 8-week-old Nicotiana mbacum L. cv. Samsun were infiltrated with bacterial suspensions. Inoculated plants were incubated in a growth chamber at 27°C with a 14/10 h daylight regime and visually monitored for reactions. For testing the prevention of the HRlike response, cells of AC5070 (105 CFU/ml) were infiltrated into tobacco leaves. The preinoculated areas were reinoculated with 2×10^4 CFU of AC5070 per ml or 5×10^6 CFU of P. syringae pv. pisi Psp1 per ml at desired intervals.

Cloning of hrpN_{Ecc} DNA and nucleotide sequence analysis.

The genomic library of E. carotovora subsp. carotovora strain Ecc71 in pLARF5 was screened by in situ colony hybridization with a 0.75-kb internal ClaI fragment of hrpN of E. chrysanthemi (Bauer et al. 1995). Two cosmids, pAKC921 and pAKC922, that hybridized with the probe were isolated. The subclones (pAKC923 and pAKC924, Table 1) carrying hrpN DNA were used for sequence analysis.

Unidirectional 5' to 3' deletions of pAKC924 were made and the overlapping deletions differing in size by approximately 200 bp were used for sequence analysis with the Sequenase System II (U.S. Biochemicals, Cleveland, OH). In addition, we used oligonucleotide primers to verify and complete the sequence of hrpN_{5cc} with pAKC923 and pAKC924 DNAs as templates. Alignment of protein sequences was performed using the Genetics Computer Group, Inc. (Madison, WI) software program at the DNA Core facility on the University of Missouri-Columbia campus and the PC/GENE program (IntelliGenetics, Inc., Mountain View, CA). The sequence of hrpN_{Ecc} has been deposited at GenBank and has been assigned accession number L78834.

Northern blot analysis.

Bacterial cultures were grown to a value of approximately 200 Klett units at 28°C in SYG medium with or without

Table L. Bacterial strains and plasmids		Reference or source
Bacteria	Relevant characteristics	
Erwinia carotovora subsp. carotovora		Zink et al. 1984
71		Muraua et al. 1991
AC5006	Lac mutant of 71	This study
	RamA*, EMS mutant of AC5006	Chamerjee et al. 1995
AC\$041	Nall derivative of AC5006	Chatterjee et al. 1995
AC5047	non A - mini-To S-Km mutant of AC3047, All 1144	Chatterjee et al. 1995
AC5070	Obligative of ACS070, Rama, Kitt, Spc	This study
AC5090 .	Ohl derivative of AC5041, RsmA*, Km	• •
AC5093		
Pseudomonas syringae pv. pist	WITH A name	A. J. Novacky
Pspl	Wild type	
•	m. misco i JD27 annà i andà i thini	BRL, Frederick, MD
Escherichia coli	980lacZ AMI5, AlacZYA-argF), UI69 hsdRI7 recAl endAl thi-1	Zink et al. 1984
DH5a	nmAl locy hsdS20(rB mB), recolo (para)	Castilho et al. 1984
HB101	* 4 P E Jack(Y7YA Werk*	Castilho et al. 1984
M8820	A TOTAL CONTINUE OF THE APPRAISMENT CANAL	Gray and Greenberg 1992
PO11734	aras(lac-proAB) rpsL \$80lacZ_sMIS recAS6	
VJS533	madan branch to the	
Plasmids	·	Chamerjee et al. 1995
pAKC852	Ohll', Tel	This study
pAKC863	Ohir, 15 Derived from pAKC852, ohll::Mudl, Kmf, Tcf	Cui et al. 1995
PAKC880	RsmA*, Spc*	This study
	RsmA*, Spc* pLARF5 containing htpNsc from genomic library of Ecc71, Tc* pLARF5 containing htpNsc from genomic library of Ecc71, Tc*	This study
pAKC921	pLARF5 containing http://ex.from genomic library of Ecc71, Te* pLARF5 containing http://ex.from genomic library of Ecc71, Te*	This study
pAKC922	pLARF5 containing http://www.mom.genome.noising.http://www.cloned.into.pSK*.Apf 4.0-kb EcoRI fragment of pAKC921 containing http://www.cloned.into.pSK*.Apf	This study
pAKC923	4.0-kb EcoRI fragment of pAKC921 containing http://ecc.cloned into pSK*. Apr 1.4-kb EcoRI fragment of pAKC922 containing http://ecc.cloned.into.psK*. Apr	Lemer and Inouye 1990
pAKC924	Spcf	Baner et al. 1995
pCL1920		Keen et al. 1988
pCPP2172	hrpN _{Ed} Ap	Keen et al. 1988
DLARF5	Te'	Figurski and Helinski 197
DRK415	Te ^c	Stratigene, La Jolla, CA
pRK2013	Mob*, Tra*, Km²	Strangers, La John, Co.
pBluescript SK+	Apr	Gray and Greenberg 1992
pHV200	8.8-kb luz DNA in pBR322, Apr	Pearson et al. 1994
PH V 200	Frameshift mutation of luxI in pHV200, Ap	ated as Hal in our previous pu

^{*} Uncommon abbreviations: EMS = ethyl methane sulfonate; Ohl = N-(3-oxohexanoyl)-L-homoserine lactone, designated as Hal in our previous publications; ramA = regulator of secondary metabolites; hrpNg = E. carotovora subsp. carotovora DNA fragment carrying a hrpNg homolog (Bauer et al. 1995).

spectinomycin. The procedures for RNA isolation and Northern blot analysis described in Chatterjee et al. (1991) and Liu et al. (1993) were followed. A 0.7-kb Acci-Smal internal fragment of hrpN_{Eee} was used as the probe.

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HARPIN IS NOT NECESSARY FOR THE PATHO-GENICITY OF ERWINIA STEWARTII ON MAIZE. Musharaf Ahmad, D. R. Majerczak, and D. L. Coplin*. Dept. of Plant Pathology, The Ohio State University, Columbus, OH 43210-1087, USA.

Erwinia stewartii elicits a hypersensitive response (HR) in tobacco if expression of the hrp-like wis regulon is enhanced. A clone containing E. amylovora hrpNE was used as a hybridization probe to locate a gene for harpin production, hrpNEs, within the wts gene cluster. Transposon mutagenesis and complementation analysis revealed that hrpNes is a monocistronic operon. Sequence analysis Indicated that it encodes a 382-amino acid; glycine-rich polypeptide, which lacks cysteine and an N-terminal signal peptide. Harpines is 58% identical and 78% homologous to harpiness and 41% identical and 66% homologous to harpinech from E. chrysanthemi. Purified harpines was professe sensitive and heat-stable, and it elicited a typical HR in tobacco leaves. Antibodies to harpin Es cross-reacted with harpings and conversely. Harpings was found in cytoplasmic, membrane, and extracellular fractions. Chromosomal mutations in hrpNEs were constructed by Tn5 mutagenesis and marker-exchange. The mutants were HRand did not produce detectable harpin in Western blots. However, they remained fully pathogenic on maize seedlings with respect to symptom severity, ED50 and response time, and they grew as well as the wild-type strain in planta. Likewise, loss of harpin did not affect the ability of a hrpNes mutant to grow endophytically in several grasses. wtsB, wtsD, and wtsF mutants accumulated Harpines intracellularly, indicating that these DNA regions are necessary for harpin secretion.

Molecular differentiation of *Erwinia amylovora* strains from North America and of two Asian pear pathogens by analyses of PFGE patterns and *hrpN* genes

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Summary

In order to determine a possible genomic divergence of Erwinia amylovora 'fruit tree' and raspberry strains from North America, several isolates were differentiated by pulsed-field gel electrophoresis (PFGE) analysis, the size of short DNA sequence repeats (SSRs) and the nucleotide and deduced amino acid sequences of their hrpN genes. By PFGE analysis European strains are highly related, whereas strains from North America were diverse and were further distinguished by the SSR numbers from plasmid pEA29. The E. amylovora strains from Europe showed identical HrpN sequences in contrast to the American isolates from fruit trees and raspberry. Those were related to each other, but distinguishable by their HrpN patterns. The Asian pear pathogens differed in HrpN among each other and from E. amylovora. Erwinia pyrifoliae isolates and the Erwinia strains from Japan were ordered via their HrpN sequences in agreement with the PFGE patterns. For all three pathogens, dendrograms from PFGE and sequence data indicate an evolutionary diversity within the species in spite of a genetic conservation for parts of the hrpN genes suggesting a long persistence of the Asian pear pathogens in Korea and Japan as well as of fire blight in North America. Some of the divergent American E. amylovora isolates share PFGE patterns with the relatively uniform European strains.

Introduction

Fire blight of apple and pear fruit trees and raspberry as well as of other rosaceaous plants is assumed to have originated in the Eastern part of North America, from

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where the disease might have been endemic for a long time, and was then distributed in the last century to many countries of the Northern hemisphere and to New Zealand (Bonn and van der Zwet, 2000). In Korea, a bacterial disease of pears and its causative agent *Erwinia pyrifoliae* has been described (Rhim et al., 1999), which was distinguished from *Erwinia amylovora* by molecular and microbiological tools (Kim et al., 1999) and additional DNA sequences (McGhee et al., 2002). Another disease, bacterial shoot blight of pear, was noticed on the island of Hokkaido in Japan (Beer et al., 1996) and the pathogen has been shown to be more related to *E. pyrifoliae* than to *E. amylovora* (Kim et al., 2001a).

Erwinia amylovora has been extensively investigated for many physiological, biochemical and molecular features (reviewed in Vanneste, 2000). Two main factors are a strict requirement for pathogenicity: the ability to produce the acidic exopolysaccharide (EPS) amylovoran, encoded in the 17 kb ams region of the chromosome (Bugert and Geider, 1995) and to induce a hypersensitive response (HR) on non-host plants, encoded by the 30 kb hrp region (Kim and Beer, 2000). The large number of hrp genes is associated with regulation and transport of two elicitor proteins, HrpN (harpin) (Wei et al., 1992) and HrpW (Barny, 1995). The adjacent dsp region with dspA/E (Gaudriault et al., 1997; Bogdanove et al., 1998) may contribute to harpin activity. Because mutagenesis of hrpN revealed residual HR-inducing activity of HrpN-fragments, HrpN might not be strictly required as an intact protein (Barny, 1995) and conservation of its sequence has not been strongly selected in mutational changes during evolution. Accordingly, its DNA and amino acid sequences could be open to changes without affecting bacterial fitness and may be useful for strain and species differentiation.

Another molecular tool for differentiation of *E. amylovora* and *E. pyrifoliae* as well as the *Erwinia* strains from Japan is PFGE analysis (Zhang and Geider, 1997; Zhang et al., 1998; Jock et al., 2002). Macrorestriction of the bacterial genome revealed several closely related but distinguishable pattern types for *E. amylovora* which were used to follow spread of fire blight in Europe and in the Mediterranean region (Jock et al., 2002). Another method to distinguish *E. amylovora* strains and the *Erwinia* strains

from Japan was determination of short sequence DNA repeats (SSR) in the PCR fragment amplified with primers P29A and P29B from the common E. amylovora plasmid pEA29 (Kim and Geider, 1999), also applied to the Erwinia strains from Japan (Jock et al., 2003a). In contrast to E. amylovora strains from Europe and the Mediterranean region, heterogeneous PFGE patterns of American strains could indicate a long persistence of the pathogen in North America. Based on HrpN-sequences, E. pyrifoliae strains from Korea (Kim et al., 2001b) and pear-pathogenic Erwinia strains from Japan (Kim et al., 2001a) were also divergent. Accordingly, macrorestriction and hrpN sequence analysis can be used for differentiation and grouping of strains within the three pathogens.

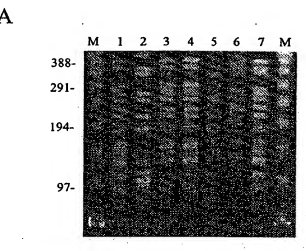
Results

PFGE patterns of E. amylovora strains isolated in North America from fruit trees and raspberry

To estimate possible diversity of Erwinia amylovora strains in North America, we collected a set of strains in several areas of Eastern Canada. The samples were derived from fire blight-infected orchards with pear and apple trees located in Nova Scotia near Kentville and in Ontario near Toronto as well as from hawthorn adjacent to the apple orchard in the Kentville area. After an Xbal digest (Fig. 1A, Table 1), the strains isolated from hawthorn and apple trees from Kentville carry the PFGE pattern Pt4 as found before (Jock et al., 2002) for strains isolated in England, Western France and Northern Spain. Strikingly, the strains from pears which are isolated in Nova Scotia in an orchard only 100 km apart from the apple orchard, had a different pattern. Another divergent pattern type was found for strains isolated in pear orchards of the Ontario region. The divergence or similarity of the investigated isolates can be deduced from the dendrogram in Fig. 1B.

An additional set of strains was isolated in Eastern Canada 1997 in the Kentville area of Nova Scotia. Strains from apple trees had the same pattern as the strains from hawthorn and apple isolated in 2000 (Table 1). Some shared the PFGE pattern with the European pattern types Pt1, others with Pt4. Most others were quite divergent in contrast to the closely related European pattern types.

Remarkably, E. amylovora strains isolated in Europe and in the Mediterranean region have an identical PFGE pattern in an Spel digest except for one band shifted for strains of the Xbal pattern type Pt3 (Zhang and Geider, 1997). In contrast, the strains from America were divergent in their Spel pattern (Fig. 2A), except strains EaCa4/ 97 and EaCa6/97 with an identical Spel pattern, which were isolated in the same year and area. Three strains which were isolated in Eastern Canada from raspberry, an alternative host for fire blight, differed in their PFGE



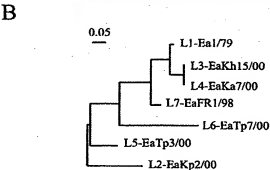


Fig. 1. PFGE analysis of E. amylovora strains isolated in Canada after genomic Xbal digests.

A. Lanes: M: λ DNA marker (sizes at left in kb); 1: Ea1/79 (Pt1, standard pattern for central Europe); 2: EaKp2/00; 3:EaKh15/00; 4: EaKa7/00; 5: EaTp3/00; 6: EaTp7/00 (Isolates from Eastern Canada.); 7: EaFR1/98 (from Germany); Xbal digests. B. Dendrogram from patterns in A. Bar, distance scale.

patterns after Xbal and Spel digests among each other and showed barely overlapping patterns with 'fruit tree' strains (Fig. 2A, Table 1). The raspberry strain IL6 from Illinois is more related to the 'fruit tree' strain Ea1/79 than the other rubus strains assayed.

The sizes of SSRs of strains from a narrow region of Eastern Canada

A more variable feature than PFGE patterns of the E. amylovora genome is a DNA fragment from the common plasmid pEA29 with several short sequence DNA repeats (Kim and Geider, 1999; Jock et al., 2003a). The SSR numbers are not related to the PFGE patterns, enabling differentiation of strains with the same pattern by SSR numbers. Rarely, the SSR numbers differ for strains isolated from plants in the same region. Nevertheless, strains

Table 1. Bacteria used in the experiments.

Strain	Description of isolation (plant, place, year, provider)	PFGE pattern
E. amylovora strains from	Canada (fruit tree)	
EaCa1/00	pear (P. communis), Annapolis Valley/Nova Scotia, 2000, G. Braun	A (d)
EaCa4/97	apple (M. domestica), Annapolis Valley/Nova Scotia, 1997, G. Braun	B/Pt4 (a)
EaCa6/97		
	apple, Annapolis Valley/Nova Scotia, 1997, G. Braun	B/Pt4 (a)
aCaH6	Harrow, D. Hunter	B/Pt4
EaCaH9	Harrow, D. Hunter	Pt1
EaCaL4	London, D. Hunter	B/Pt4
EaCaS16	Simcoe, D. Hunter	Pl1
aCaS23	Simcoe, D. Hunter	Pt1
aCaS5	Simcoe, D. Hunter	Pt1
aCaV15	Niagara, D. Hunter	₿y
EaCaV18	Niagara, D. Hunter	Bz
EaCaV8	Niagara, D. Hunter	Bx
aCaW2E	Wentowth country/Hamilton, D. Hunter	Pt1
aCaW3	Wentowth country/Hamilton, D. Hunter	B/Pt4
EaKa6/00	apple (M. domestica), Kentville, this work	B/Pt4
EaKa7/00	apple, Kentville/Nova Scotia, 2000, this work	B/Pt4
EaKa8/00	apple, Kentville/Nova Scotia, 2000, this work	. -
aKa9/00	apple, Kentville/Nova Scotla, 2000, this work	_
aKa10/00	apple, Kentville/Nova Scotia, 2000, this work	· _
aKh14/00	hawthorn (Crataegus sp.), Kentville/Nova Scotla, 2000, this work	
aKh15/00	hawthorn (Crataegus sp.), Kentville/Nova Scotia, 2000, this work	B/Pt4
aKh17/00	hawthorn, Kentville/Nova Scolia, 2000, this work	B/Pt4
aKp1/00	pear (<i>P. communis</i>), Kentville/Nova Scotia, 2000, this work	A
aKp2/00	pear (<i>P. communis</i>), Kentville/Nova Scotia, 2000, this work	Α
aKp5/00	pear (P. communis), Kentville/Nova Scotia, 2000, this work	_
aTp3/00	pear (P. communis), Niagara Falls/Ontario, 2000, this work	C
аТр7/00	pear (<i>P. communis</i>), Niagara Falls/Ontario, 2000, this work	D
aTp9/00	pear (<i>P. communis</i>), Niagara Falls/Ontario, 2000, this work	. -
aTp10/00	pear (<i>P. communis</i>), Niagara Falls/Ontario, 2000. this work	_
aTp12/00	pear (<i>P. communis</i>), Niagara Falls/Ontario, 2000, this work	-
aTpyr6/00	Asian pear (P. pyrifolia), Niagara Falls/Ontario, 2000, this work	
E. amylovora strains from CA1R	apple, California, A. Jones	E
CA263	apple or pear, California, A. Jones	E
CASR	apple, California, A. Jones	F
a88	pear, Washington, A. Jones	E
a110	apple, Michigan, A. Jones	B/Pt4
a153	USA, L. Pusey	
		Pt1
aU8/96	apple, Utah, 1996 (Bereswill <i>et al.</i> , 1998)	-
L01	A. Jones	B/Pt4
B93-5	pear, Idaho, A. Jones	E
13-1	Indian hawthorn, Louislana, A. Jones	G ·
_1196	pear, Washington, A. Jones	Ē
A029	pear, Washington, A. Jones	Ē
A033	pear, Washington, A. Jones	E
P100 ·	apple, Washington, A. Jones	E
R1	pear, Oregon, A. Jones	Ε
R6	pear, Oregon, A. Jones	Ē
TRJ2	apple, Utah, A. Jones	B/Pt4
/SDA14	apple, Washington, A. Jones	B/Pt4
SDA34	apple, Washington, A. Jones	E
amulnunra etraine from	raspberry (isolated in North America)	
aCa1/95		/- \
	raspberry (Rubus ideus), Annapolis Valley, Nova Scotia	– (b)
aCa1/98	raspberry, Bouctouche, New Brunswick	– (bx)
aCa8/96	raspberry, Bouctouche, New Brunswick	- (c)
MR1	raspberry, Michigan	K`
aRKK3	raspberry, Michigan	Ĵ
aRUB7	raspberry (Bereswill et al., 1998)	ĭ
		•
<u>6</u>	raspberry, Illinois	H (e)
amulawara etraine from	Europe (Jock et al., 2002)	
	Crataegus sp., France, JP. Paulin	Pt3a
FBP1430	0 1 1 0 1000	
FBP1430 a1/79	Cotoneaster sp., Germany, 1979	Pt1 (a)
FBP1430 a1/79	Cotoneaster sp., Germany, 1979 P. communis, Toulouse (France), 1994	Pt1 (a) Pt4
FBP1430 a1/79 a9–7	P. communis, Toulouse (France), 1994	Pt4
a1/79 a1/79 a9–7 a296 a321		

Table 1. Cont.

Strain	Description of isolation (plant, place, year, provider)		PFGE pattem
EaUK2/98	hawthorn, Kent (UK), 1998		Pt1
P1573	Cotoneaster sp., Dorset (UK), 1995, A. Aspin	•	Pt4
E. pvrifoliae strains fro	m Korea (Kim et al., 2001b)		¥00
Ep1/96	Asian pear (Pyrus pyrifolia), South Korea, 1996		· PtA
Ep4/97	Asian pear (P. pyrifolia) South Korea, 1996		PtB
Ep28/96	Asian pear (P. pyrifolia), South Korea, 1996		. PIC
Ep31/96	Asian pear (P. pyrifolia), South Korea, 1996	•	PtC
Ep102/98	Asian pear (P. pyrifolia), South Korea, 1998		PtA
Erwinia strains from Ja	span (Kim <i>et al.</i> , 2001a)		•
Eip546	Asian pear (P. pyrifolia), Hokkaido, 1979, A. Tanii		other
Eip547°	Aslan pear (P. pyrifolia), Hokkaido, 1979, A. Tanil		PtJp1
Ejp556	Asian pear (P. pyrifolia), Hokkaido, 1994, A. Tanii		other
Ejp557	Asian pear (P. pyrifolia), Hokkaido, 1994, A. Tanii		PtJp1
Ejp562	Asian pear (P. pyrifolia), Hokkaido, 1994, A. Tanil	•	PUp1
Ejp617	Asian pear (P. pyrifolia), Hokkaido, 1996, R. Roberts		other

a. Letters A to K refer to the pattern of Xbal digests, as for Pt1 to Pt4 and PtJp1; highly related pattern are listed with Y, similar patterns with a lower case letter added to the main type in upper case. (a to e) in this column refer to Spel-digests of genomic DNA as for PtA, PtB and PtC of E. pyrifoliae. -, not assayed.

from a narrow area in Nova Scotia were not identical in SSRs displaying numbers of 5, 7, or 9 (Fig. 3, Table 2). These data suggest independent changes of E. amylovora populations for SSR. In particular, a strain (EaTp12/ 00) isolated from a pear tree in the neighbourhood of the orchard, where other strains listed in Table 2, had been isolated, showed a divergent SSR number.

Sequence analysis of the hrpN genes of E. amylovora 'fruit tree' and raspberry strains

The hrpN genes from several E. amylovora 'fruit tree' strains with divergent PFGE patterns and from three raspberry strains were cloned by PCR amplification. The European 'fruit tree' strains Ea1/79, CFBP1430, Ea321 (nucleotide sequence from data library), Ea9-3, P1573 or EaFR3/97 with pattern Pt1, Pt3 (2x), Pt4 or Pt1A, respectively, showed almost identical nucleotide sequences for their hrpN genes with differences of not more than one nucleotide. On the other hand, the American raspberry strains could be distinguished by their HrpN sequences from 'fruit tree' strains from North America. Three motifs in the N-terminal part are typical for rubus strains and can even be considered diagnostic for their distinction from 'fruit tree' strains (Fig. 3A, boxes). In addition, the rubus strain EaCA1/95 showed a six amino acid insertion sequence in the centre of HrpN and a smaller insertion closer to the N-terminus. These sequences distinguished strain EaCA1/95 from strains EaMR1 and IL6 (Fig. 3A, underlined). In a dendrogram, the 'fruit tree' strain Ea1/79 from Germany is well separated from the aligned American rubus strains, but all E. amylovora strains differ in their

Table 2. SSR numbers of E. amylovora strains isolated 2000 in Eastern Canada.

Origin	Isolated from	Name	SSR
Kentville	pear	EaKp1/00	7
•	• •	EaKp2/00	7
		EaKp5/00	7
	apple	EaKa6/00	9
		EaKa7/00	9
		EaKa8/00	8
		EaKa9/00	7
	•	EaKa10/00	5
	hawthorn	EaKh14/00	>10
		EaKh15/00	8
	•	EaKh17/00	8
Toronto	P. pyrifolia	EaTpyr6/00	4
	pear	EaTp9/00	4
	•	EaTp10/00	4
	pear*	EaTp12/00	3

a. From tree adjacent to main orchard.

alignment patterns from the Asian pear pathogens (Fig. 3B).

Sequence analysis of the hrpN genes of E. pyrifoliae strains and Erwinia strains from Japan

Erwinia amylovora 'fruit tree' and raspberry strains share motifs of HrpN with the Asian pear pathogens. In Fig. 3A, the sequences of the Korean Erwinia pyrifoliae Ep1/96 and of an Erwinia strain from Japan, Ejp557, were aligned for their possible relationship to the E. amylovora raspberry strains. Erwinia pyrifoliae strains and the Erwinia strains from Japan were strikingly distinct from both E.

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b. Previously named Ejp546a, derived from a culture obtained with Ejp546.

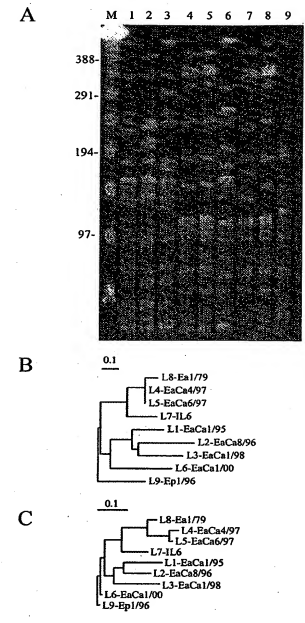


Fig. 2. PFGE analysis of *E. amylovora* strains isolated from raspberry in Canada and Illinois by genomic *Spet* digests in comparison with isolates from apple in Canada and an *E. amylovora* isolate from cotoneaster and an *E. pyrifoliae* strain.

A. Lanes: M: λ DNA marker (sizes at left in kb); 1: EaCa1/95 (rb); 2: EaCa8/96 (rb); 3: EaCa1/98 (rb); 4: EaCa4/97 (a); 5: EaCa6/97 (a); 6: EaCa1/00 (p); 7: IL6 (rb); 8: Ea1/79 (highest band from partial digest); 9: Ep1/96 (*E. pyrifoliae*).

B. Dendrogram from pattern in A.

C. Dendrogram from pattern of Xbal digest with the strains applied in A. Suffix 'a', isolated from apple; 'p', from pear; 'IL6', from raspberry. Bars, distance scales.

amylovora groups. The HrpN sequences of the two Asian pear pathogens were related to each other, but not identical and differed in at least four clusters of more than two amino acids.

The *E. pyrifoliae* strains Ep1/96 and Ep102/98 belong to the PFGE pattern type PtA, Ep4/97 to PtB and Ep28/96, Ep31/96 to pattern type PtC (Kim *et al.*, 2001b). Most parts of their HrpN sequences were identical. Nevertheless, Ep1/96, Ep4/97 and Ep102/98 showed a DNA insertion encoding seven amino acids, which distinguished them from the others (Fig. 4). The motif 'GGSGGGL' is reiterated twice for these strains, but is not conserved for *E. amylovora* or the *Erwinia* strains from Japan (Fig. 3A and Fig. 4A). The distance scale in the dendrogram derived in Fig. 4B indicates a close relationship of the investigated *E. pyrifoliae* strains with small differences. Ep1/96, Ep28/96 and Ep102/98 are highly related, less Ep31/96, whereas Ep4/97 is more distinct from the others.

The *Erwinia* strains from Japan analysed were also not completely homogenous for their HrpN sequences. Strains Ejp547, Ejp557 and Ejp562 were highly related in the PFGE patterns after *Xbal* digests, whereas the others could be separated from the first group on this basis (Kim *et al.*, 2001a). In agreement with those data, the HrpN proteins of Ejp547, Ejp557 and Ejp562 differed from the amino acid sequences derived from the other strains at five sites of HrpN (Fig. 5A). The dendrogram in Fig. 5B indicates the relationship of Ejp557, Ejp547 and Ejp562, separating them from the other strains, also confirming PFGE data that strain Ejp547 is not identical with strain Ejp546, obtained in the same agar culture.

The sequences of the hrpN genes of E. amylovora fruit tree' and raspberry strains as well as of E. pyrifoliae strains and Erwinia strains from Japan showed a differential degree of conservation. The E. amylovora 'fruit tree' and rubus strains were 97% related to each other, whereas the HrpN proteins of E. pyrifoliae and Erwinia strains from Japan had only 83% similarity to HrpN of E. amylovora 'fruit tree' strains.

A summarizing dendrogram (Fig. 6) grouped the *E. amylovora* strains apart from the two Asian pear pathogens. *E. pyrifoliae* strains are highly related to each other, and less to the *Erwinia* strains from Japan.

Discussion

The PFGE patterns of the strains isolated in North America are divergent, in contrast to the pattern of strains from Central Europe and the Mediterranean region, which were grouped into four main pattern types (Jock et al., 2002). In spite of basically unrestricted trade in fruit and fire blight host plants, there has been no obvious mixing of pattern types in Europe and the Mediterranean region. Sequential spread from infected sites is the dominant way of disease

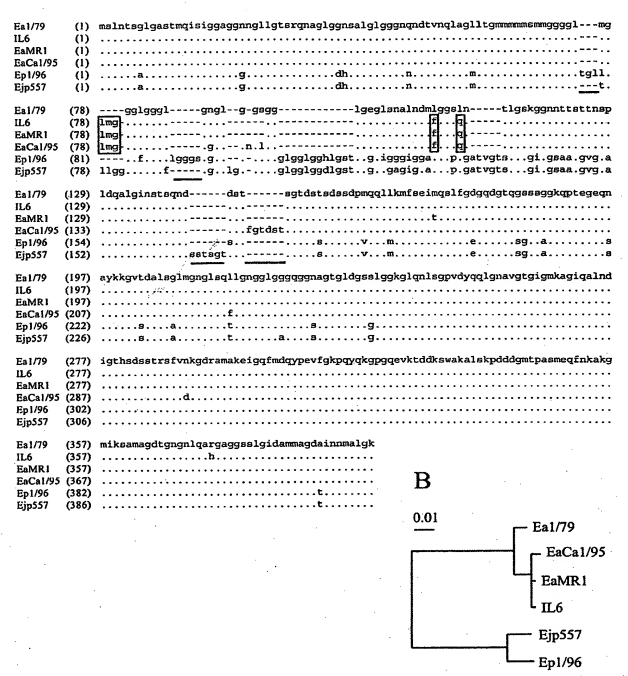


Fig. 3. Sequence alignment of the HrpN proteins from raspberry strains isolated in Canada compared with the E. amylovora 'fruit tree strain' Ea1/79, E. pyrifoliae Ep1/96 and Ejp557, an Erwinia strain from Japan. A. Common motifs for raspberry strains are boxed and unique insertions for strains are underlined. B. Dendrogram from the amino acid sequences aligned in A. Bar, distance scale.

A

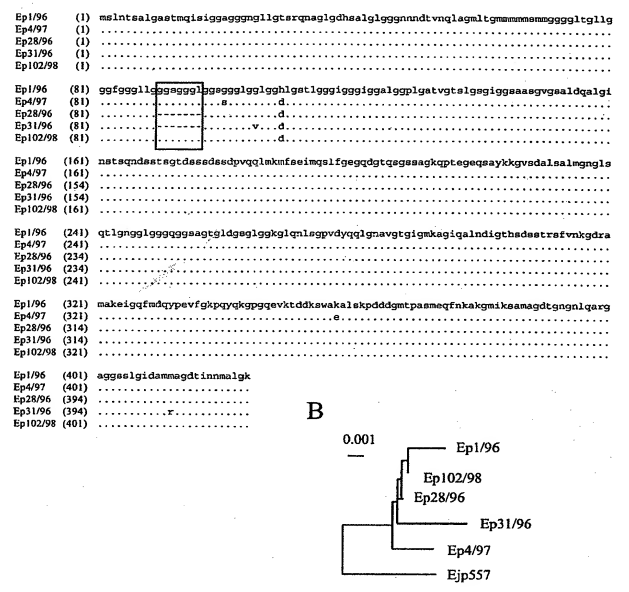


Fig. 4. Comparison of HrpN proteins from five *E. pyrifolia* strains.

A. Amino acid aligment. The motif for strain differentiation is boxed.

B. Dendrogram from the amino acid sequences aligned in A. Bar, distance scale.

distribution, except for introduction of fire blight into Central Spain and Northern Italy, where plant imports can be connected with appearance of fire blight caused by *E. amylovora* strains displaying pattern type Pt3, which has not been found in the adjacent regions.

An ordered PFGE pattern was not found for strains from North America, because even a relatively low number of

isolates gave rise to several different patterns. They also differ from European patterns except for Pt1 and Pt4, which were found repeatedly in isolates from Eastern Canada. Thus, Pt1 and Pt4 could have originated in North America and were then distributed to Europe (Jock et al., 2002), first to England with the first European fire blight outbreaks (Billing and Berrie, 2002). The other patterns in

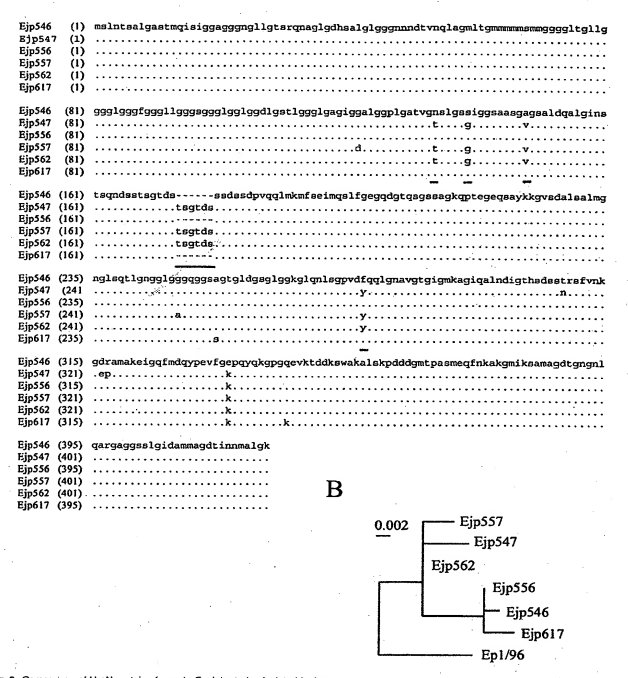


Fig. 5. Comparison of HrpN proteins from six Erwinia strains isolated in Japan. A. Amino acid alignment. An insertion motif and amino acid substitutions for strains Ejp547, Ejp557, and Ejp562 are underlined. B. Dendrogram from the amino acid sequences aligned in A. Bar, distance scale.

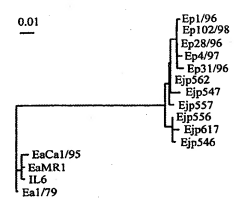


Fig. 6. A dendrogram showing the relatedness of *E. amylovora* 'fruit tree' and raspberry strains and the evolutionary distance of the Asian pear pathogens based on the HrpN amino acid sequences. Bar, distance scale.

America can be explained by genetic changes over a long time period; they were rarely distributed to other countries except for Pt2, a pattern which was found first in Egypt and also in a strain from California (Zhang and Geider, 1997; Jock et al., 2002).

A special subgroup of *E. amylovora* strains from raspberry is endemic in North America and has only been isolated there. A reason for the difference in the PFGE patterns from *E. amylovora* 'fruit tree' strains could be the unusual host, which required many genomic changes for adaptation of the pathogen. On the other hand, their presumably long persistence in North America could have allowed accumulation of many base changes in the genome causing their pattern heterogeneity.

The SSR numbers are not related to PFGE patterns or the areas of isolation as also found for American (Schnabel and Jones, 1998) and European *E. amylovora* strains (Kim and Geider, 1999). Nevertheless, different numbers indicate non-identical isolates from fire blighted plants. Among intermediate numbers there is a high occurrence of low numbers such as 3 and 4, which are not often observed in Central Europe. In isolations from the same apple orchard in Kentville, we found SSR numbers from 5, 7 and 9. Normally, only one SSR-type is usually isolated in the same set of isolates, but recently, we observed some exceptions like in England where we found SSR type 3 and 7 in isolates from adjacent plants (Jock *et al.*, 2003a).

The ability to induce a hypersensitive response (HR) on non-host plants is a common feature of plant pathogenic bacteria. In evolution, many genes of the *hrp* cluster especially those involved in protein secretion have been highly conserved among bacteria (Van Gijsegem *et al.*, 1993; Bogdanove *et al.*, 1996). A spontaneous base change in *hrpL* within an *E. pyrifoliae* population has been recently described (Jock *et al.*, 2003b). Genes encoding harpins

are highly divergent even for related bacteria. The HrpN proteins of two related species such as P. stewartii ssp. stewartii (E. stewartii) and P. stewartii pv. gypsophylae show only 60% similarity to each other (EMBL Nucleotide Sequence Database accession numbers AF282857 and AF21176 respectively). The similarity of these harpins and HrpN of E. carotovora ssp. carotovora (AF302656) to harpin of E. amylovora is 62%, 56%, and 49% respectively. The sequence information of hrpN is not only suited for classification of bacterial species, but also for grouping of strains within a species. On the other hand, HrpN can be conserved, found for E. amylovora 'fruit tree' strains, where the sequences matched at the nucleotide level. These strains isolated from raspberry in North America. share more than 95% similarity. A high relationship was also observed between E. pyrifoliae strains from Korea and the Japanese pear pathogen, whereas E. amylovora strains match with these pathogens less than 85%. Although the Erwinia stains from Japan have not been taxonomically classified, the relatedness of HrpN proteins adds to the notion to place these with E. pyrifoliae into the same species (Kim et al., 2001a). In addition, HrpN sequences provided also information for strain differentiation within a species.

Because the transport of harpin depends on several cellular proteins, its sequence cannot freely change only to conserve its elicitor activity. Whether the HrpN protein or even the DspA/E-protein (Gaudriault et al., 1997; Bogdanove et al., 1998) contribute to host plant specificity of a pathogen has still to be shown. The divergences of the HrpN sequences should indicate an evolutionary drift, similar to the PFGE patterns analysed. The most likely explanation is the long persistence of E. amylovora in North America, of E. pyrifoliae in Korea and the slightly different pear pathogen in Japan. Furthermore, the occurrence of European pattern types Pt1 and Pt4 among the divergent American PFGE patterns may indicate a rare escape of fire blight from its origin in North America.

Experimental procedures

Bacterial strains, PCR and PFGE analyses

The *E. amylovora* strains were isolated in the Eastern part of Canada, or were gifts from colleagues (Table 1). They were confirmed as *E. amylovora* on several agar plates including MM2Cu (Bereswill *et al.*, 1998) and by PCR assays (Bereswill *et al.*, 1992). Pulsed-field gel electrophoresis analysis (Jock *et al.*, 2002) and determination of the SSR numbers (Kim *et al.*, 1999) were done as described. *Erwinia pyrifoliae* (Kim *et al.*, 2001b) and the *Erwinia* strains from Japan were also described previously (Kim *et al.*, 2001a). For pattern comparison, the PFGE fragments were assigned by eye with letters and the program CLUSTALX1.81 used for pairwise alignments. The dendrograms were adjusted with NJ-tree and further processed in a graphics program. Pattern analy-

sis was also done with the public domain programs ImageJ (v. 1.30; W. Rasband, NIH, USA) and Cross Checker (v. 2.91; J. B. Buntjer, Wageningen, the Netherlands) and alignment with Treecon vs. 1.3b (Y. van de Peer, Konstanz, Germany) and CLUSTALX1.81 respectively. Corrections by eye were required for further adjustment of the band assignments.

Analysis of the hrpN genes from E. amylovora and the Asian pear pathogens

The hrpN genes of strains from the three pathogens were amplified with PCR consensus primers, which were deduced by comparison of several known nucleotide sequences from plant pathogens namely E. amylovora (EMBL Nucleotide Sequence Database accession number M92994) or P. stewartii (accession number AF282857). Primer HRPN1 was 5'-ATGAGTCTGAATACAAG-3' (at start of E. amylovora hrpN) and primer HRPN3c 5'-GCTTGCCAAGTGCCATA-3' hrpN, 11 bp downstream from stop codon). In some cases, weak PCR bands obtained could indicate incomplete matching of the primers. The amplified DNA fragments were cloned into vector pGEM-T and were commercially sequenced. To cover the total hrpN genes, a third primer HRPMc (5'-CCACGGCGTTACCCAACTGCTGG-3') located in the central part of the hrpN gene was used to cover gaps in the HrpN sequences. Alignments and dendrograms were created with

Erwinia pyrifoliae and the Erwinia strains from Japan were considered to be sufficiently related to E. amylovora to amplify their hrpN genes with the Erwinia PCR consensus primers given above. This was indeed possible and allowed cloning and sequencing of their hrpN DNA fragments as for E. amylovora by using primers HRPN1 and HrpN3c. A primer comprising the stop codon at the C-terminus of hrpN did not result in the formation of a PCR product together with primer HRPN1.

The hrpN nucleotide sequences from strains Ea1/79, EaCa1/95, IL6, EaMR1, Ejp546, Ejp557, Ep1/96, Ep31/96, Ep4/97 and were deposited in the EMBL Nucleotide Sequence Database with the accession numbers AJ579689 (Ea1/79), AJ579690 (EaCa1/95), AJ579691 (IL6), AJ579692 (EaMR1), AJ579693 (Ejp546), AJ579694 (Ejp557), AJ579695 (Ep1/96), AJ579696 (Ep31/96) and AJ579697 (Ep4/97).

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The HrpZ Proteins of *Pseudomonas syringae* pvs. *syringae*, *glycinea*, and *tomato* Are Encoded by an Operon Containing *Yersinia ysc* Homologs and Elicit the Hypersensitive Response in Tomato but not Soybean

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The Pseudomonas syringae pathovars are composed of host-specific plant pathogens that characteristically elicit the defense-associated hypersensitive response (HR) in nonhost plants. P. s. pv. syringae 61 secretes an HR elicitor, harpiness (HrpZPss), in a hrp-dependent manner. An internal fragment of the P. s. pv. syringae 61 hrpZ gene was used to clone the hrpZ locus from P. s. pv. glycinea race 4 (bacterial blight of soybean) and P. s. pv. tomato DC3000 (bacterial speck of tomato). DNA sequence analysis revealed that hrpZ is the second ORF in a polycistronic operon. The amino acid sequence identities of HrpZ_{Ps}/HrpZ_{Psc} and HrpZ_{Ps}/HrpZ_{Psi} were 79 and 63%. respectively. Although none of the HrpZ proteins showed significant overall sequence similarity with other known proteins, HrpZ_{Pst} contained a 24-amino acid sequence that is homologous with a region of the PopA1 elicitor protein of the tomato pathogen, Pscudomonas solanacearum GMI1000. hrpA, the upstream ORF, was highly divergent: The amino acid sequence identities of HrpAps, HrpAps and HrpA_{Ps}/HrpA_{Pst} were 91 and 28%, respectively, and no HrpA sequence showed similarity to known proteins. In contrast, the predicted products of the downstream ORFs in P. s. pv. syringae and P. s. pv. tomato, hrpB. hrpC, hrpD, and hrpE showed varying levels of similarity to those of yscl. yscl. ysck, and yscl. These are colinearly arranged genes in the virC locus of Iersinia spp., which are involved in the secretion of the Yop virulence proteins via the type III pathway. The similarity of the Ysc proteins was generally stronger in comparisons with the P. s. pv. tomato Hrp proteins. The HrpZ proteins were purified by heat denaturation of contaminating problems followed by ammonium sulfate fractionation, hydrophobic chromatography, and gel electrophoresis. All three HrpZ proteins elicited the HR in tomato, whereas none of them elicited significant necrosis in soybean. The results indicate that HrpZ is encoded in an operon containing some of the genes involved in its own secretion and suggest that HrpZ structure does not directly determine bacterial host range.

Phytopathogenic strains of Pseudomonas syringae cause two patterns of necrosis when the bacteria invade a plant. On a susceptible ("compatible") host, a necrotic lesion often develops over a period of days, with necrosis spreading as the bacteria multiply and the plant becomes diseased. On a resistant or nonhost plant, a localized cellular necrosis is induced within 24 to 48 h. and bacterial multiplication is inhibited. This was first reported by Klement (1963; Klement et al. 1964), who observed that when high concentrations of pathogenic bacteria are infiltrated into an incompatible plant they elicit a visible necrosis which is limited to the infiltrated area. This reaction, called the hypersensitive response (HR), involves localized cell death and production of anti-microbial compounds at the site of pathogen invasion (Bonas 1994). The ability of P. syringae and other nontumorigenic, gramnegative, bacterial pathogens to elicit the HR is governed by hrp genes. Typical Hrp mutants are pleiotropically defective in planta: They do not elicit the HR in nonhosts and they fail to multiply and cause disease in host plants (Lindgren et al. 1986). Clusters of hrp genes have been identified in many gram-negative phytopathogenic bacteria (Bonas 1994). A 25kh hrp cluster from P. s. pv. syringae 61 is sufficient to confer the tobacco HR phenotype, but not the pathogenic phenotype on nonpathogenic bacteria (Huang et al. 1988). hrp genes have also been cloned and characterized extensively from P. s. pv. phaseolicola NPS3121. P. solanacearum GM1000. Xanthomonas campestris pv. vesicatoria 75-3, and Erwinia amylovora Ea321 (Lindgren et al. 1986; Boucher et al. 1987; Beer et al. 1991; Bonas et al. 1991). Certain hrp genes are widely conserved among these pathogens, and several encode components of a protein secretion pathway that is similar to the type III pathway used by Yersinia, Shigella, and Salmonella spp. to secrete extracellular proteins involved in animal pathogenesis (Van Gijsegem et al. 1993). One activity of the hrp-encoded secretion pathway in phytopathogenic bacteria is the secretion of proteinaceous elicitors of the HR, which are also encoded by hrp genes.

The first hrp-encoded elicitor characterized was harping, from E. amylovora (Wei et al. 1992). Similar elicitors have since been isolated from other bacteria, including P. s. pv. syringae 61, P. solanacearum GMI1000, and E. chrysanthenii

EC16 (He et al. 1993; Arlat et al. 1994; Bauer et al. 1994). Proteins in this family of elicitors share several general characteristics. They are glycine rich, heat-stable, lack cysteine, and appear highly susceptible to proteolysis. They lack an Nterminal signal peptide, but they are secreted to the bacterial milieu. Their expression and secretion is dependent on hrp genes. The biological role of these proteins in pathogenesis has not yet been determined, but the purified proteins can induce an HR on a nonhost plant such as tobacco. However, there are significant differences in the organization of the elicitor operons and the activity of the elicitors, which suggests that the Erwinia harpins, the P. syringae hrpZ product and the P. solanacearum popA product may represent three distinct classes of elicitors. In this work we will refer to the P. s. pv. syringae elicitor as HrpZPss tather than harpinpss (He et al. 1993). This distinction is supported by the weak similarity of the amino acid sequences of the four proteins, with the only exception being the C-terminal halves of the Erwinia harpins (Bauer et al. 1994).

The location of known elicitor genes in reference to the hrp cluster varies in P. s. pv. syringae, P. solanacearum, and E. amylovora, hrpN and hrpZ are contiguous or within the hrp cluster, whereas popA lies outside (although near) the P. solanacearum hrp cluster (Wei et al. 1992; He et al. 1993; Arlat et al. 1994). There are no genes downstream of the elicitor gene in either the hrpN or the popA operons, which means that mutations in the elicitor genes do not have a polar effect on the Hrp phenotype, and mutant construction is straightforward. In contrast, mutagenesis and complementation studies of the hrp cluster from P. s. pv. syringae 61 have indicated that hrpZ lies upstream of at least one other hrp gene within an operon (Huang et al. 1991; Xiao et al. 1992).

In E. amylovora and E. chrysanthemi, harpins have been demonstrated to be sufficient and necessary to elicit the HR, and mutation of hrpN in E. amylovora has shown that harpine is required for pathogenesis (Wei et al. 1992). However hrpN mutants of E. chrysanthemi can establish infections, albeit at a significantly reduced frequency, which suggests that harpine, is important but not essential for pathogenesis (Bauer et al. 1995). In contrast, a popA mutant of P. solanacearum is fully pathogenic on susceptible hosts, indicating that PopA1 is not required for pathogenesis (Arlat et al. 1994).

These elicitors may play a role in controlling the host specificity exhibited by E. amylovora and plant pathogenic pseudomonads such as P. syringae and P. solanacearum. However it is difficult to compare the activity of HrpZ_{Pss} and harpine, in host and nonhost plants because legumes and rosaceous plants, the hosts of P. s. pv. syringae 61 and E. amylovora Ea321, respectively, respond poorly to preparations of any of these elicitor proteins (Wei et al. 1992; He et al. 1993). PopA1 from P. solanacearum does appear to act in a hostspecific manner, inducing an HR on resistant lines of petunia and the nonhost tobacco, but not on susceptible lines of petunia or tomato (Arlat et al. 1994). This phenotype is similar to that of avr genes, but PopA1 is distinct from known Avr proteins in eliciting the HR directly on resistant plants. Harpinech elicits an HR on some compatible hosts of E. chrysanthemi, but in contrast to the other three bacteria E. chrysanthemi is a broad-host range pathogen and the activity of harpinech may not be representative of elicitor activity in a highly host-specific system (Bauer et al. 1995).

In previous work we cloned and characterized the hrpZ gene from P. s. pv. svringae 61, a weak pathogen of bean, and demonstrated with Southern and immunoblots that other pathovars of P. syringae contain homologs of this gene (He et al. 1993). This supported the hypothesis that HrpZ represents a family of elicitors common to all pathogenic strains of P. syringae. We report here the isolation of homologs of HrpZPss from two other experimentally important pathovars of P. syringae-P. s. pv. tomato and P. s. pv. glycinea. Examining HrpZ from these three pathovars enabled us to look within this family of elicitors for variations in sequence and activity which could indicate a role in host range determination. In addition, we characterized the two genes flanking hrpZ in P. s. pv. syringae and P. s. pv. glycinea and the entire hrpZ operon of P. s. pv. tomato. In conjunction with an accompanying paper (Huang et al. 1995), this completes the sequence of the P. s. pv. syringae 61 hrp genes carried on pHIR11 and provides clues to the function of the genes downstream of hrpZ. A preliminary account of portions of this work has been published (Collmer et al. 1994).

RESULTS

Cloning hrpZ from P. s. pv. tomato and P. s. pv. glycinea.

We previously used Southern hybridization to demonstrate that both P. s. pv. glycinea race 4 and P. s. pv tomato DC3000 contain sequences homologous to a 0.75 kb BstXI internal fragment of hrpZ from P. s. pv. syringae (He et al. 1993). The same probe was used to screen genomic libraries of P. s. pv. glycinea and P. s. pv. tomato. The libraries were constructed in E. coli DH5a by inserting 8- to 12-kb fragments from partial Sau3AI digests of genomic DNA into the BamHI site of pUCP19. The screen identified two plasmids with inserts of approximately 10 kb: pCPP2201 (P. s. pv. tomato) and pCPP2200 (P. s. pv. glycinea). The same BstXI fragment was used to probe a Southern blot of pCPP2201 and pCPP2200 digested with BamHI, EcoRI, and PstI. The probe identified two PsiI fragments of 2.2 and 2.4 kb from pCPP2201 and pCPP2200 respectively (Fig. 1). The two Pstl fragments were cloned into the PstI site of pBluescript II SK(-) (Stratagene, La Jolla, CA) in E. coli DH5a to create the plasmids pCPP2202 to pCPP2205, with the inserts in both orientations with respect to the lac promoter. Cell lysates of E. coli DH5α containing pCPP2203 (hrpZ_{Psi} in the vector promoter orientation) and pCPP2202 (hrpZ_{Psg} in the vector promoter orientation) induced an HR on tobacco, but those from cells containing pCPP2205 (hrpZ_{Pn} in the opposite orientation of the vector promoter) and pCPP2204 (hrpZ_{Pss} in the opposite orientation of the vector promoter) did not. HR activity was retained after incubating the lysate for 10 min at 100°C and removing denatured proteins by centrifugation. Insensitivity to heat treatment is a characteristic feature of previously isolated HR elicitors. Proteins in the lysates were separated on an SDS-polyacrylamide gel, transferred to an Immobilon-P membrane and immunoblotted with antibodies raised against purified HrpZ_{Pss}. Cross-reacting proteins of a similar size to HrpZ_{Pss} were observed and provisionally named HrpZ_{Psg} and HrpZ_{Psi} (Fig. 2, lanes 2 and 4).

The intensity of the $HrpZ_{Psg}$ and $HrpZ_{Pss}$ bands was quite low in comparison to the band for $HrpZ_{Pss}$ expressed from pSYH10 in *E. coli* DH5 α (Fig. 2, lane 1). This implied either

that expression was low due to the distance of the cloned gene from the *lac* promoter or that HrpZ_{Psg} and HrpZ_{Psg} did not hybridize strongly to the antibodies. A band corresponding to HrpZ_{Psg} from pSYH10 could be clearly seen on a Coomassie-stained gel, but the bands for HrpZ_{Psg} and HrpZ_{Psg} were indistinct, which implies that low expression was a primary reason for the low signal. In an attempt to improve the level of expression of HrpZ_{Psg} and HrpZ_{Psg} we subcloned *Eco*RI-BamHI fragments containing the inserts from pCPP2202 and pCPP2203 behind the T7 promoter of pET21(+) in *E. coli* BL21(DE3) to create the plasmids pCPP2206 and pCPP2207.

The T7 promoter enabled a moderate improvement in protein expression (Fig. 2, lanes 3 and 5).

A common arrangement of ORFs in the hrpZ operons of P. s. pv. syringae, P. s. pv. glycinea, and P. s. pv. tomato revealed by DNA sequence analysis.

Previously, we determined the complete nucleotide sequence of hrpZ from P. s. pv. syringae by sequencing a 1.4-kb subclone of pHIR11 (a cosmid containing the entire hrp cluster from P. s. pv. syringae) (He et al. 1993). In addition, analysis of the complementation groups and transcriptional

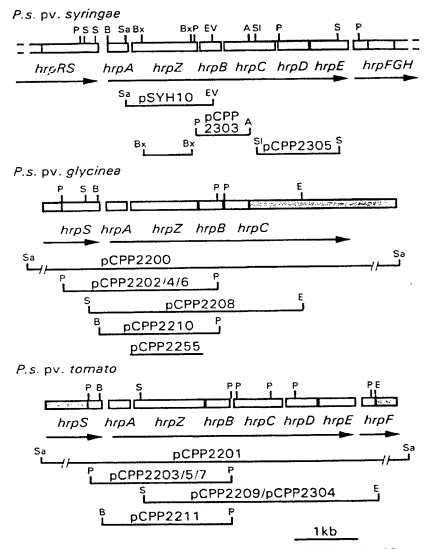


Fig. 1. Physical maps of the hrpZ regions from Pseudomanas syringae pv. syringae 61, P. s. pv. glycinea tace 4, and P. s. pv. tomato DC3000 and clone-used in this study. Open boxes represent sequenced ORFs: filled boxes represent unsequenced DNA. Direction of transcription is indicated by the arrows. Key restriction sites within the sequenced regions are indicated, along with the subclones used in this study. The 0.75-kb BstXI fragment from $hrpZ_{pi}$ used as a probe for hrpZ genes in other pathovars is also shown. Restriction endonuclease abbreviations: A, $Agel^*$; B, Bglli; Bx, $BstXII^*$; E, EcoRl, EV, $EcoRV^*$, P, Pstl, S, Sucl. Sa, $Sau3A^*$; Sl, $Sal1^*$, *Not all sites are shown

units of pHIR11 using TnphoA and Tn5-gusA1 mutagenesis (Huang et al. 1991: Xiao et al. 1992) suggested that hrpZ lay within an operon, upstream of at least one other hrp gene. Further subclones of pHIR11 were used to determine the sequence of the entire hrpZ_{Pss} operon (this study. Huang et al. 1995). We also determined the sequence of (i) the 2.2- and 2.4-kb PsrI subclones from pCPP2201 (hrpZ_{Psr}⁺) and pCPP2200 (hrpZ_{Psg}*), (ii) an overlapping 3.7-kb SacI-EcoRI subclone from pCPP2201 (designated pCPP2209), and (iii) part of an overlapping 3.6-kb subclone from pCPP2200 (designated pCPP2208), as shown in Figure 1. This yielded the sequence of the entire P. s. pv. tomato hrpZ operon and the first half of the P. s. pv. glycinea operon. The sequenced region of P. s. pv. syringae and P. s. pv. tomato extends from hrpS (Xiao et al. 1994), through the hrpZ operon to the beginning of the hrpH operon (Huang et al. 1992), demonstrating that the organization of this region of the hrp cluster is conserved in both pathovars.

Codon preference analysis of the DNA sequence, using *P. s. pv. syringae* codon usage data, predicted that *hrpZ* was the second of six ORFs, all oriented in the same direction, an arrangement conserved in *P. s. pv. tomato* and at least the first four ORFs of *P. s. pv. glycinea*. The sequence of the noncoding DNA is shown in Figure 3. Five of the six ORFs have clear potential ribosome binding sites. The fifth ORF has a putative ribosome binding site in *P. s. pv. syringae*, but the site in *P. s. pv. tomato* is less clear, the initiation codon shown being selected by alignment with the ORF in *P. s. pv. syringae*. In the absence of recognizable terminator elements downstream of the first five ORFs it seems likely that the six ORFs represent a single operon, transcribed from upstream of the first ORF. The five predicted ORFs were provisionally named *hrpA* through *hrpE*, as shown in Figures 1 and 3.

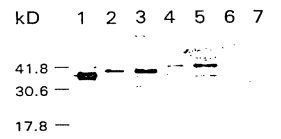


Fig. 2. Immunobiot showing expression of cloned httpZ in E. coli. Cultures were grown in LM to an OD₂₀₀ of 0.8 to 1.0 at 30°C, collected by centrifugation and resuspended in 5 mM MES, pH 5.5. For lanes 3, 5 and 7, and 4, T7 expression was induced with 1 mM IPTG when the cells reached an OD₂₀₀ of 0.6. 3 h prior to collection. The cells were disrupted by sonication, and the crude lysate was partially purified by removal of the insoin-fee fraction after incubating the samples at 100°C for 10 min. SDS-loading buffer was added and the samples were incubated at 100°C for 2 min. The proteins were resolved by SDS-polyacrylamide gel electrophoresis. Following electrophoresis the proteins were transferred to Immobilon-P membrane (Millipore, Bedford, MA), probed with anti-HtpZe₂₀ antibodies and visualized with goat anti-rabbit antibody conjugated with alkaline phosphatase. Lanes: 1, E. coli. DH5α (pSYH10) (HtpZe₂₀₁); 2, E. coli. DH5α (pCPP2203)(HtpZe₂₀₂); 5, E. coli. BL21(DE3) (pCPP2207)(HtpZe₂₀₁); 6, E. coli. DH5α (pBluescript II); 7, E. coli. BL21(DE3) (pCPP2207)(HtpZe₂₀₁); 6, E. coli. DH5α (pBluescript III); 7, E. coli. BL21(DE3) (pCPP2207)(HtpZe₂₀₁); 7, E. coli.

A hrp/avr promoter consensus sequence lies upstream of the hrpZ operons of the three P. syringae pathovars.

The conserved sequence GGAACC-16bp-CCACNNA lies 50 bp upstream of the initiation codon of hrpA in all three pathovars (Fig. 3). This motif has been identified in the promoter regions of many avr and hrp genes (Innes et al. 1993; Shen and Keen 1993), and appears to be involved in positive regulation by HrpL, a putative alternative sigma factor which is itself positively regulated by HrpR and HrpS (Xiao and Hutcheson 1994). HrpL is a member of a family of alternative sigma factors, many of which are involved in secretion of extracellular factors in response to environmental stimuli (Lonetto et al. 1992). The presence of this promoter motif further supports the suggestion that the six ORFs form a single transcriptional unit which is regulated in a hrp-dependent manner. This motif can also be found beyond hrpE, upstream of hrpFGH in P. s. pv. syringae and P. s. pv. tomato, as indicated at the bottom of Figure 3, suggesting that the latter three ORFs form an independent hrp-regulated transcriptional unit in these two pathovars.

Comparison of the HrpZ proteins of the three P. syringae pathovars.

The predicted amino acid sequences for HrpZ from each of the three pathovars are aligned in Figure 4. Although the proteins migrate slightly anomalously on an SDS polyacrylamide gel, the relative sizes of the estimated molecular weights correspond to the predicted values, with $HrpZ_{Pst}$ being the largest of the three proteins (36.5 kDa), followed by $HrpZ_{Psg}$ (35.3 kDa) and $HrpZ_{Pss}$ (34.7 kDa). Amino-terminal sequencing of the first 10 to 15 residues of purified $HrpZ_{Psg}$ and $HrpZ_{Ps}$ confirmed the predicted initiation codons of both proteins, which aligned with the start codon of HrpZ_{Pss} as shown in Figures 3 and 4. The proteins expressed in E. coli appear to be the same size as those recovered from the supernatants of P. s. pv. glycinea and P. s. pv. tomato, indicating that the cloned gene is intact and that there are no large posttranslational modifications or deletions of HrpZ taking place in P. syringae but not in E. coli.

The amino acid sequence of HrpZ_{Pst} is quite highly conserved with respect to HrpZ_{Pst}, having 87% similarity and 79% identity. HrpZ_{Pst} is less conserved with respect to the two other proteins, with 75% similarity and 63% identity to HrpZ_{Pst}. However, the physical features of HrpZ_{Pst} and HrpZ_{Pst} are almost identical to those reported for HrpZ_{Pst} (He et al. 1993). All three are glycine-rich proteins lacking cysteine and tyrosine. HrpZ_{Pst} is the most glycine rich, being 15.7% glycine. The proteins lack the hydrophobic signal sequence used to target proteins for secretion via the Sec export pathway (Pugsley 1989). Analysis of the amino acid sequence fails to identify any obviously significant secondary structure, which is consistent with their sensitivity to proteases, and supports the suggestion that they adopt a fairly open structure in aqueous solution.

In our previous analysis of HrpZ_{Pss} (He et al. 1993), we noted the presence of two sets of short, direct repeats. Only one of these repeats. GGGLGTP, is conserved in the three proteins, with the substitution of a serine for threonine in the first repeat of both HrpZ_{Psg} and HrpZ_{Pss}. The significance of these repeats, if any, is unknown. A database search with each of the three proteins using the BLAST algorithm (Altschul et

syringae	TTTTTTGCAG	AAGATCTGGA	ACCGATTCGC	GGACACATGC	CACCTAGCTG
glycinea	TITITIGCA.	GAGCGCTGG	L ACCGATTTAL	L CCCTCCTTN C	CAGMA mome
tomato	TTTTTTGCAA	AGACGCTGG	ACCGTATCGC	AGGCTGCTGC	CACTACTCAC
syringae	TACCAAGCAA	TTACGCTGGT	CACAGACGAAG	GGGTATGACG	max ma
glycinea	TACCAAGCAA	TTACGCTGGT) ACACACCAAC	GGGTATCACG	TIATG
tomato	TACCADGCAD	TOTOCOTOC	ACAGACC <u>AAC</u>	GGGTATCACG	TTATG
comaco	TACCAAGCAA	1CACGC16G1	AAAICITAAG	GGGCATCAAA	TCATG
			,.		5000013464
syringae		321bp	<u>1</u>	GATTTCTTG.	ACGCCCCTTC
glycinea	hrpA	321bp		GATTTCTTGA	ATGCCCCCAT
tomato		336bp		AATTATTTCT	CATTOCCCCCAI
			ā	Service C	
syringae	ATACCTGAGG	GGGCTGCTAC	* TTTTACCACC	TTGTG .ATG	=====
glycinea	CACACAGAGG	CCCCTCCTAC	TTTTAGGAGG	TIGIG. AIG	
	DOADACACACAC	GGGCIGCIAC	TTTGAGGAGG	TTGTG . ATG	
tomato	TCATCAGAGG	GGGCCGCTAC	CTTGGGATGG	GCGTTTTATG	
	=> <===	====		· · · ·	
syringae		1020bp			
glycinea	hrpZ	1032bp			
tomato		1107bn			
		770, ND			
syringae					
glycinea					
tomato					
syringae	TGACCGACAA	CCGCCTGACG	GAGAACTCAC	GTG	
glycinea	TGACTGATAC	CCGCCTGACG	GAGLECTORC	GTG	
tomato	TGACTGLCLG	CCGCCTGACG	CACLACCACE	G1G	nrps
	201.010.10.10.10	CCGCC1GACG	GHGHACCAG.	GTG	
ermina.	3 C O imm				•
syringae	36300		TAGAGGTTTC	CGTG	
glycinea	36 ADD		T <u>AGAGG</u> TTCT	CGTG	
tomato	3€9bp		TAGAGGTTTC	CGTG	
syringae		dd108			
glycinea	hrpC	incomplete			
tomato		801bb			

syringae			CT CCECT COC		
tomato		TGA13	GACCIGACCG	CCGAGGACTA	TTGGACTCAG
LOa		ATGATG	AGCCTTTCTG	CCGAGGATCA	CTGGATTCAC
syringae	TGGTGGTGCA	ATCCCTGGCC	ATGGGCGCAT	CCGGGGCTGGC	AAAGCCGGTT
tomato	TGGTGGTGCA	ACCCCTGGCA	GTGGGCACAT	TCGGAGTGGC	ATGLCCGLTT
syringae	CGCCGAGCGC	TGCGGACTGA	CCGTCAGCGZ	ATGTGLAGOO	בית מידים -
tomato	CGCCAACGCT	CGTGGGGTTLT	CCCTCTCTCT	CTOCOL TOCOL	CITAIG
		corcerin.	CGGICAGIGA	C: C	CIUMIG
syringae		h	26.61		
		nrpu	386pb		
tomato			396bp		
syringae	TGAGTAT.	.CCGCTCCTC	TCTGCACCAG	GAATTOTOCO	ATG
tomato	TGARTCCG.	AACCAGCTTC	TCTGCATCAG	GAATACGCCC	ATC
			<u>- : - 0</u>		1410
syringae	hrpE	576hn			
tomato		576bb			TGA
	_	J . U.D			TGA
syringae	Abozerom	_	mmaaaa		
	AACAGACT	C	TIGUGGCGAA	AATGGAACCG	CTCCACCTGT
tomato	TACACACTOT (UTGCACTCAC	TTGATCGCAT	GATGGAACCG	CTCGGCGGGT
syringae	TIGOTCCACT (CAAGGTTTGA	ACCTTTCTGC	TGGAGTATCL	GGACATG
tomato	TTGCTCCACT (CAAGGTTTGL	ACCCTTCTGC	TGGAGCACCA	CCICITO
				AUJAJOAGOL	GGALAIG

Fig. 3. Nucleotide sequences of the noncoding regions of the httpZ operon from Pseudomonas syringae pv. syringae, P. s. pv. glycinea, and P. s. pv. tonato. The sequences flanking the six ORFs of the httpZ operon were aligned using the PILEUP algorithm (Genetics Computer Group). For P. s. pv. syringae and P. s. pv. tonato the sequence extends from immediately downstream of httpS to the end of the operon. For P. s. pv. glycinea the sequenced region terminates at the beginning of httpC. The proposed initiation and termination codons are highlighted for each ORF. The http/avr consensus sequences upstream of httpA amd httpF are marked by double lines, with the conserved nucleotides in bold and the putative ribosome binding sites for each ORF underlined. A short inverted repeat upstream of http2 is also indicated with dashed arrows.

al. 1990) did not find significant homology to any other bacterial proteins, with the exception of a single, glycine rich region found only in $HrpZ_{Psi}$ (Fig. 4). This stretch of 24 amino acids has homology at both the nucleotide and amino acid level to a region of the host-specific elicitor PopA1 from P.

solanacearum, as shown at the bottom of Figure 4. There is no overall similarity of the amino acid and nucleotide sequences of HrpZ to the HR elicitors characterized from E. amylovora, E. chrysanthemi, and P. solanacearum except to a degree accounted for by their similar composition.

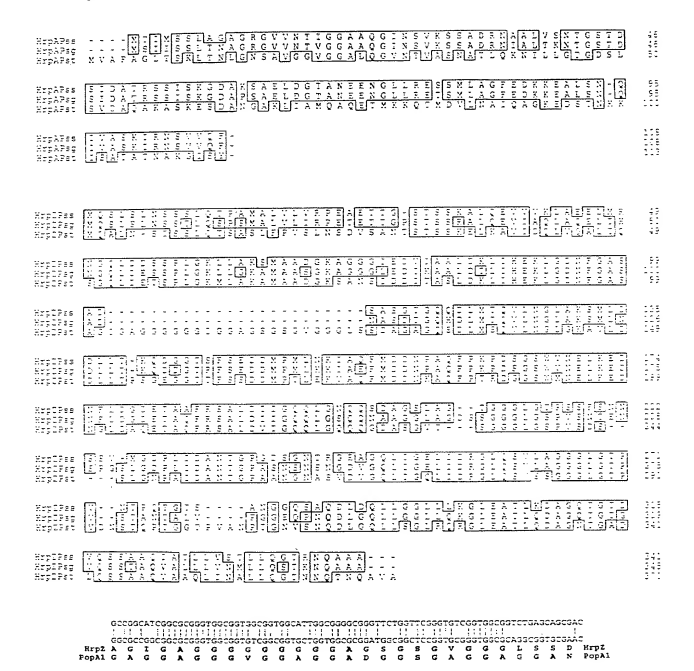


Fig. 4. Alignment of the protein sequences of HrpA and HrpZ. The predicted protein sequences of HrpA and HrpZ from Pseudomonas syringae pv. syringae, P. s. pv. glycinea, and P. s. pv. tomato were aligned using the PILEUP algorithm (Genetics Computer Group). The alignment of a unique glycine rich region of HrpZ_{Pst} with a homologous region of PopA1 from P. solanacearum is also shown.

The predicted HrpA protein of *P. s.* pv. tomato differs substantially from that of *P. s.* pv. syringae and *P. s.* pv. glycinea.

The first ORF of the hrpZ operon starts 50 bp downstream of the conserved hrp/avr promoter motif, as shown in Figure 3. The predicted product is a small (11 kDa), hydrophilic protein with a hydrophobic N-terminus. An alignment of the amino acid sequences from all three pathovars is shown in Figure 4. Although the predicted sequences of HrpA from P. s. pv. syringae and P. s. pv. glycinea are highly conserved, with 92% similarity and 91% identity to each other. HrpA from P. s. pv. tomato is quite divergent, having only 42% similarity and 28% identity to HrpA from P. s. pv. syringae The presence of a ribosome binding site and the highly conserved character of HrpA in two of the three pathovars supports the hypothesis that HrpA is translated. T7 polymerasedependent expression of hrpA (described below) provides further evidence for production of a HrpA protein. Cell lysates of E. coli expressing only HrpA did not elicit the HR on tobacco (data not shown), which suggests that it does not contribute directly to the HR. The role of HrpA in the bacterium is unknown, and it shows no significant homology to any previously characterized proteins.

T7 expression studies.

To confirm the production of proteins corresponding to the two sets of newly cloned hrpA and hrpZ genes, the BgIII-PstI fragments from P. s. pv. glycinea and P. s. pv. tomato were subcloned into pET21(+) and the products specifically labelled by T7 promoter/polymerase-dependent expression in E. coli BL21(DE3) cells incubated with [35]-methionine (Studier et al. 1990). Radiolabeled proteins in the cell lysate were analyzed by SDS-polyacrylamide gel electrophoresis and autoradiography (Fig. 5). Lysates of cells containing pCPP2211 displayed unique bands which corresponded well with the predicted molecular weight of HrpA (11.5 kDa) and were consistent with the previously observed mobility of HrpZ_{PR} (Fig. 5, lane 2). Lysates of cells containing pCPP2210 contained bands corresponding to HrpZ_{Fsg} (36 kDa) and HrpA (11 kDa)(Fig. 5, lane 3). No HrpB band was visible in the products of pCPP2211 (Fig. 5, lane 2), but this could potentially be attributed to the omission of cysteine, which is not required for HrpA and HrpZ synthesis, from the amino-acids added to the reaction mixture. T7 expression of HrpB was independently confirmed for both P. s. pv. syringae and P. s. pv. tomato using a 0.84-kb PstI-AgeI fragment of pHIR11 and the 3.7-kb SacI-EcoRI fragment from pCPP2209, subcloned into LITMUS 28 to construct the plasmids pCPP2303 and pCPP2304. T7 expression in E. coli BL21(DE3) cells was performed as outlined above and in Figure 5. In each case a protein of about 13 kDa was observed, which corresponds well with the predicted molecular weight of HrpB from each of the two pathovars (data not shown). In an accompanying study Huang et al. (1995) have confirmed the production of proteins corresponding to HrpC. HrpD. and HrpE from P. s. pv. syringae 61. The similarities between the three pathovars suggest that the equivalent ORFs in P. s. pv. glycinea and P. s. pv. tomato also encode proteins. However when we independently confirmed the production of HrpD from $P_{\rm c}$ s. pv. syringae 61 using a 1.3-kb Sall-Sacl subclone from pHIR11 cloned into pT7-6 (pCPP2305) our results suggested the use of an alternative initiation codon to make a larger (21 kDa) HrpD protein (data not shown). In the absence of a strong ribosome binding site at either of the putative initiation codons, the exact size of HrpD remains uncertain.

The four ORFs downstream of hrpZ show varying similarities to Yersinia Ysc proteins.

The hrpC, hrpD, and hrpE genes downstream of hrpZ in P. s. pv. syringae 61 have been sequenced and the products identified using T7 polymerase-dependent expression (Huang et al. 1995). Two of the predicted proteins, HrpC and HrpE, were shown to be homologous to the proteins YscJ and YscL, respectively, which are encoded in the virC operon of Yersinia enterocolitica and are involved in the type III secretion pathway (Michiels et al. 1991). Homologs of YscJ have also been found in the hrp clusters of several other phytopathogenic bacteria, including P. solanacearum and X. campestris (Fenselau et al. 1992: Gough et al. 1992). Additional homologs are Salmonella typhimurium FliF and Rhizobium fredii NolT (Jones et al. 1989; Meinhardt et al. 1993). The same four downstream ORFs are found in P. s. pv. tomato. and the partial sequence of the operon from P. s. pv. glycinea confirms the presence of the first two of these ORFs, hrpB and hrpC, in this pathovar (Fig. 6).

HrpB is fairly conserved in all three pathovars, as shown by the alignment presented in Figure 6. It encodes a small serine-rich protein of approximately 13 kDa. BLAST searches using HrpB from either P. s. pv. syringae or P. s. pv. glycinea identified no significant homologies, but a search using HrpB from P. s. pv. tomato identified similarity to the Yersinia protein, Yscl. Yscl is 115 amino acids long, thus slightly shorter than HrpB (127 amino acids). yscl lies immediately upstream of yscJ in the virC operon, which suggests that the downstream ORFs of the hrpZ operon might be colinear with a region of the virC operon.

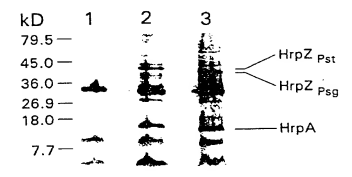


Fig. 5. T7 polymerase-dependent expression and radiolabeling of HrpA and HrpZ. T7 promoter/polymerase expression was carried out using the pET21(+) v ctor system in *E. coli* BL21(DE3). Cells were grown in LM to an OD_{NR} of 0.5, then centrifuged and resuspended in M9 minimal medium supplemented with 0.01% amino acids (lacking methionine and cysteine), glucose and thiamine. Cells were incubated at 30°C for 3 h and then induced with 1 mM IPTG for 10 min, followed by incubation with nfampicin at 300 μ g/ml for 30 min. Cells were incubated with 10 μ Ci [18 S]-methionine for 10 min, lysed in SDS-loading buffer, and the proteins were separated by SDS-polyacrylamide electrophoresis and visualized by autoradiography. *E. coli* BL21(DE3) cells carried the following plasmids in lanes: 1, pET21(+); 2, pCPP2211; 3, pCPP2210.

The apparent colinear arrangement of this group of hrp and ysc genes led us to inspect the P. s. pv. syringae and P. s. pv. tomato HrpD proteins for possible similarity to the Yersinia spp. YscK proteins. The similarity between the HrpD of P. s. pv. syringae and Y. pseudotuberculosis was the highest, with 28% of the amino acids identical and 57% similar. The HrpD and YscK proteins are of similar overall composition, and they lack any predicted transmembrane segments. However, there is a striking discrepancy between the sizes of the two proteins. HrpD is only 133 amino-acids long, whereas YscK from Y. pseudotuberculosis is 209 amino-acids long. From the T7 experiments described above it is important to note that in the absence of a strong ribosome binding site, the precise ini-

tiation codon of the hrpD ORF is uncertain; it is conceivable that hrpD actually initiates immediately downstream of hrpC, at the ATG codon which overlaps the stop codon of hrpC, which would yield a predicted protein of 176 amino acids for HrpZ_{Pst} or 175 amino acids for HrpZ_{Pst} in an arrangement similar to that of the yscJ and yscK ORFs in Yersinia spp. However, this codon and all other potential initiation codons upstream of the one we have chosen lack ribosome binding sites, and the pattern of codon usage suggests that the intergenic region is not translated.

Although the similarities between HrpB/YscI, HrpD/YscK. and HrpE/YscL are lower than those involving HrpC/YscJ. the similarities of HrpB/YscI and HrpE/YscL are clearly in-

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Fig. 6. Alignment of the protein sequences of HrpB from Pseudomonas syringae pv. syringae, P. s. pv. glycinea, and P. s. pv. tomato, and HrpC, HrpD and HrpE from P. s. pv. syringae and P. s. pv. tomato with Yscl. Yscl. Ysck, and Yscl. from Y. enterocolitica and Y. pseudotuberculosis (Michiels et al. 1991; Rimpilainen et al. 1992). (continued on next page)

dicative of probable homology as based on a difference between the scores for the optimized and the average of 100 random Gap alignments being at least 5 times the standard deviation for the randomized alignments (Doolittle 1986). The scores for HrpD/YscK lie at the margin of significance by this measure. However, the varying levels of similarity are consistent with the divergence observed between Hrp proteins from different *P. syringae* pathovars and between Ysc proteins from different *Yersinia* spp. The results for HrpB.C and E lend support to the weak homology of HrpD to YscK and suggest that hrpB, hrpC, hrpD, and hrpE are colinear with yscl, yscJ, yscK, and yscL.

In a recent report. Van Gijsegem et al. (1995) observe that the *P. solanacearum* GMI1000 *hrp* cluster also encodes homologs of YscJ and YscL but not YscI and YscK. It is possible that with relatively divergent Hrp sequences, similarities with Ysc proteins may be found only after examining the sequences from several plant pathogens. It is interesting to note that there is no ORF following *hrpE* that is homologous to the protein encoded by the final gene of the *virC* operon, YscM. However, the *hrpZ* operon lies immediately upstream of the *hrpH* operon (Fig. 1), and HrpH is a homolog of YscC, a secretion protein which lies upstream of *yscIJKL* within the *virC* operon (Michiels et al. 1991). This suggests that a sig-

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Fig. 6. (continued from preceding page)

nificant proportion of the virC operon is conserved in P. syringae, albeit in a rearranged form. Eckhardt (1978) gels of total DNA, Southern-blotted and probed with a 0.75-kb BstXI internal fragment of hrpZ_{Pss}, suggested that the hrp genes are chromosomal in the three strains of P. syringae studied, rather than being plasmid-borne as are the hrp genes of P. solanacearum GMI1000 or the vsc genes of Yersinia spp. (Van Gijsegem et al. 1993, data not shown). The homologies of the hrpZ operons are summarized in Table 1.

Overexpression, purification, and biological assay of HrpZpst and HrpZpsg.

Partially purified lysates of E. coli expressing HrpZPs1 and HrpZ_{Ptr} elicited a clear HR on tobacco while control lysates of E. coli containing vector alone did not. However the activity of the cell lysates on the two host plants was more ambiguous. Soybean is generally unreactive to cell lysates from either pathogen, while tomato is quite sensitive and sometimes weakly reactive not only to cell lysates of E. coli expressing HrpZ, but also to control lysates of E. coli containing vector alone. To accurately evaluate the biological properties of HrpZ from each of the two pathovars, it was necessary to purify HrpZ. It was also necessary to ascertain that the HR observed on tobacco was due solely to HrpZ and not to the products of either of the two flanking ORFs. HrpA and HrpB, since HrpA and a fusion protein of HrpB were being expressed in addition to HrpZ by the original hrpZPsg and hrpZ_{Pst} clones.

As a first step towards purifying HrpZ, we attempted to increase the level of expression. From the sequence of the Pstl clones encoding hrpZ it was clear that long stretches of DNA encoding hrpA and the 3' end of hrpS (1.144 bp in $hrpZ_{Ps_E}$ pCPP2202 and 809 bp in $hrpZ_{Ps_E}$ pCPP2203) separated hrpZfrom the lac promoter in pBluescript II. A series of deletions of the 5' end of the hrpZPs clone were constructed using the Erase-a-Base system (Promega), bringing the lac promoter within 100 bp of the hrpZ initiation codon, and removing hrpA. Although cell lysates expressing the deleted clones retained HR eliciting activity, they did not show a substantial increase in gene expression. Searching for an explanation for this behavior we identified a number of potential contributing factors. The first possibility was the presence of a cis-acting sequence contained in the 100 bp remaining upstream of hrpZ_{Pst}. Using a terminator analysis program we identified a 9-bp inverted repeat located between hrpA and hrpZ (Fig. 3). Although this repeat lacks the AT-rich sequence downstream which is characteristic of many terminators, it is possible that its presence encourages premature transcription termination. Similar repeats, albeit with weaker secondary structure, can be found upstream of hrpZ_{Pss} and hrpZ_{Pss}. A second factor contributing specifically to the low expression of hrpZ_{Pst} may be the absence of a strong ribosome binding site. Finally, there could be factors related to the proteins themselves, such as a lack of stability.

To eliminate possible cis-acting sequences and to obtain clones of hrpZ_{Pst} and hrpZ_{Psg} that lack hrpA and hrpB, the hrpZ genes from both pathovars were amplified by PCR, directionally cloned into pBluescript II and transformed into E. coli DH5\alpha F'lacl. We obtained significantly increased expression of HrpZ_{Ps} using the plasmid pCPP2255 (Fig. 7), but unexpectedly, overexpression of HrpZ_{Psi} appeared to be deleterious to the cells, and plasmids recovered from transformants often showed rearrangements. To maximize expression of HrpZ_{Pst} under these conditions, we introduced subclones containing the gene behind the T7 promoter of pET21(+) (Novagen, Madison, WI). Unlike the lac promoter, the T7 promoter is less sensitive to distance effects, and expression of HrpZ_{Pst} in E. coli BL21(DE3), with pET21(+) as the vector, resulted in increased expression as shown in Figures 2 and 8. Expression in BL21(DE3) also allowed us to retain almost complete repression of hrpZ until induction with IPTG. Good expression of HrpZ_{Fst} was achieved using the plasmid pCPP2211 in E. coli BL21(DE3).

The quality of the samples obtained following partial purification of the lysates by heat treatment was quite variable. To ensure removal of the majority of the contaminating proteins and to obtain a more concentrated sample of protein, we further purified HrpZ by ammonium sulphate precipitation and hydrophobic chromatography, which as indicated in Figure 8. yielded a distinct band on a Coomassie-stained gel. Purified. active HrpZ could then be obtained by electroelution from excised gel slices. This procedure was also used to isolate

Table 1. Homologies of Pseudomonas syringae pv. syringae hrpZ operon proteins with proteins from other P. syringae pathovars and Yersinia spp.

P. s. pv. syringae	HrpA (108)*	HrpZ (341)	HrpB (124)	HrpC (268)	HrpD (133) ^d	HrpE (193)
P. s. pv. glycinea P. s. pv. tomato	(108) 91/92* (108) 28/42	(345) 79/87 (370) 63/75	(124) 94/96 (124) 68/80	(268) 90/95	(133) 78/87	(193) 76/87
Y, enterocolítica	2014-	53.15	Ysc1 (115) 22/45 24/45	YscJ (244) 35/59 38/60	YseK (203) 26/53 22/48	YscL (223) 21/47 22/46
Y. pseudotuberculosis			(115) 22/45 21/44	(244) 35/59 38/60	(209) 28/57 23/49	(221) 21/47 22/46

Number of amino acids in the protein is given in parentheses.

Percent identical and similar amino acids in comparison with the P. s. pv. syringae protein.

The first pair of values are the percent identical and similar amino acids in comparison with the P. s. pv. syringue protein; the second are in comparison

The data presented here are for the shorter of the two potential ORFs encoding httpD. The larger versions of the HttpD proteins of P. s. pv. syringae and P. s. pv. tomato would be respectively 175 and 176 amino acids long with 74/84% identity/similarity to each other.

HrpZ from the supernatants of P. s. pv. tomato and P. s. pv glycinea grown in hrp-inducing minimal media (Fig. 9). Preparations of the purified HrpZ proteins from P. s. pvs. syringae, glycinea, and tomato, at a concentration of ≥20 µM in MES buffer, were infiltrated into the leaves of tobacco, soybean, and tomato. The three proteins elicited a collapse involving >50% of the infiltrated tissue in tobacco and tomato leaves that developed within 18 h and was typical of the HR elicited by incompatible P. syringae strains, but they caused no visible reaction in soybean. It is worth noting that tobacco and tomato plants vary substantially in their sensitivity to harpin preparations. For example, some leaves on sensitive tomato plants will respond to 2 to 5 µM HrpZ_{Pst}, but ≥20 µM is required for consistent results. Furthermore, unlike tobacco, tomato plants that have responded hypersensitively to a HrpZ preparation do not respond to subsequent infiltrations of the elicitor. The spurious necroses sometimes observed were deduced to result from mechanical damage incurred during infiltration or the infiltration of preparations contaminated with salts or containing high concentrations of vector control E. coli lysates. These necroses developed much more quickly (within 4 to 6 h), and were much weaker and patchier than the confluent HR elicited by HrpZ. The fact that the HR induced by HrpZ in tomato and tobacco is an active response of host tissue was confirmed by coinfiltration of either sodium vanadate at $5^{-5} \times 10^{-5}$ M or lanthanum chloride at 1×10 M. Each of these two inhibitors of plant metabolism completely inhibited the HR elicited by HrpZ preparations from each of the three pathovars but not the necrosis caused by the other factors mentioned.

DISCUSSION

We have used the P, s, pv, syringae 61 hrpZ gene to isolate the hrpZ locus from P, s, pv, glycinea race 4 and P, s, pv, to-mato DC3000. Characterization of the hrpZ genes, products, and flanking DNA of these three pathovars has revealed the structure of the hrpZ operon, the relative variation among

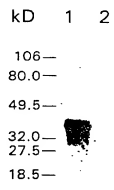


Fig. 7. Overexpression of HrpZ_{PiF} in E. coli DH5α F'lacf². Cultures were grown overnight at 30°C in LM with 1 mM IPTG. Cell lysates were partially purified by heat treatment, separated on an SDS-polyacrylamide gel, transferred to Immo² in P. immunoblotted with anti-HrpZ_{PiS} antibodies, and visualized wire goat anti-rabbit antibody conjugated with alkaline phosphatase. Lanes 1, E. coli DH5α F'lacf² ipCPP2255 ii 2 E. coli DH5α F'lacf² (pBluescript II).

ORFs within the operon, the presence of genes downstream of hrpZ that are colinear with a block of genes involved with Yersinia virulence protein secretion, and the presence in HrpZ_{Psi} of a sequence related to a sequence in the PopA1



Fig. 8. Overexpression and purification of $HrpZ_{PG}$. Cultures were grown to an OD_{GP} , of 0.6 and induced with 1 mM IPTG. $HrpZ_{PG}$ was then partially purified from the cell lysate in a three-step process: first, by heat-treatment at $100^{\circ}C$ as previously described, then by precipitation with ammonium sulphate at 30 to 45% saturation, and finally by binding to a hydrophobic resin (phenyl-sepharose) at 30% ammonium sulphate. A, Coomassie stained SDS-polyacrylamide gel. Lanes: 1, E. coli BL21(DE3)(pCP2211). B, Immoniumlolot of the samples shown in A, probed with anti- $HrpZ_{PS}$ antibodies and visualized with goat anti-rabbit antibody conjugated with alkaline phosphatase.

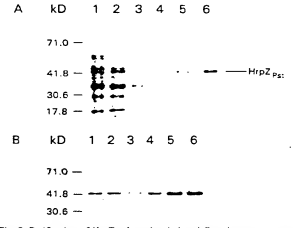


Fig. 9. Purification of HrpZ_{Pst} from hrp-induced Pseudomonas syringae pv. tomato. Cells were grown in King's broth (KB) at 30°C and then resuspended in hrp-inducing minimal medium (Huynh et al. 1989) and incubated at room temperature overnight. Cells were removed by centrifugation and the supernatant heat-treated at 100°C for 10 min. Proteins in the supernatant were precipitated with ammonium sulphate at the percent saturations indicated. Proteins were desalted, concentrated, and resuspended in 5 mM MES using Centricon-10 tubes (Amicon). A, Coomassie stained SDS-polyacrylamide gel. Lanes: 1, supernatant extracted with Strataclean resin (Stratagene). 2, heat-treated supernatant extracted with Strataclean resin (Stratagene): 3, 0 to 20% ammonium sulphate fraction: 4, 20 to 30% ammonium sulphate: 5, 30 to 40% ammonium sulphate: 6, 30 to 45% ammonium sulphate. B, Immunoblot of the samples shown in A, probed with anti-HrpZ_{Pss} antibodies and visualized with goat anti-rabbit antibody conjugated with alkaline phosphatase.

protein of the tomato pathogen P solanacearum GMI1000. We also observed that purified $HrpZ_{Pst}$ was at least as effective as $HrpZ_{Pss}$ and $HrpZ_{Psg}$ in eliciting an HR-like necrosis in the leaves of tomato, a host of P s. pv. tomato DC3000, whereas none of the HrpZ preparations elicited significant necrosis in soybean, the host of P s. pv. glycinea.

The HrpZ proteins of three P. syringae pathovars.

A comparison of the sequences of the three HrpZ proteins with each other and with HR elicitors characterized from other bacteria indicates that the HrpZ proteins represent a distinct family of elicitors that is conserved among P. syringae pathovars. The amino acid sequences of the three proteins are sufficiently similar to reveal their relatedness, but (with the exception of a sequence within HrpZ_{Pst}), they show no significant relatedness to elicitor proteins from other bacteria. Interestingly, hrpZ is the second most divergent ORF in the hrpZ operons of P. s. pv. syringae and P. s. pv. tomato, with only 63% of the predicted amino acids being identical. Nevertheless, $\text{Hrp}Z_{Pss}, \text{Hrp}Z_{Psg},$ and $\text{Hrp}Z_{Pss}$ are indistinguishable in several biological and physical properties. They have the same effect on different plants (discussed below), and they are heat stable, glycine-rich, and devoid of cysteine and tyrosine. The lack of tyrosine is a feature they differentially share with the P. solanacearum PopAl protein but not the Erwinia harpins. This property has been speculated to allow the protein to avoid the H2O2-mediated cross-linking of tyrosine residues that may occur in plant cell walls during defense responses (Bradley et al. 1992; He et al. 1993).

Interestingly, a 24 amino acid, glycine-rich stretch of HrpZ_{Psi} shows homology to part of PopA1, as does the cognate nucleotide sequence. The region of homology between HrpZ_{Psi} and PopA1 corresponds exactly to the insertion in HrpZ_{Psi}. The insertion of this element within HrpZ_{Psi} sequences that are otherwise similar among the three HrpZ proteins suggests horizontal transfer and a common ancestry with PopA1. Because the host range of *P. solanacearum* overlaps with that of *P. s. pv. tomato*, it is tempting to speculate that this region has some particular significance to pathogenesis on tomato, although, as discussed below, this is not obvious from the different effects of the two proteins on tomato.

The presence of this insert in active HrpZ_{Pst} is another indicator of the apparent plasticity of structure/function relationships in these glycine-rich elicitor proteins. That significant changes to the structure of these proteins does not abolish their activity was previously demonstrated when a fortuitous hrpZ_{Pst} clone was found to produce an active derivative of HrpZ missing the N-terminal 125 amino acids, and the popA product was observed to be degraded in culture to an active form missing the N-terminal 93 amino acids (He et al. 1993; Arlat et al. 1994). Clearly the presence of this "additional" internal sequence does not diminish the ability of the protein to elicit the HR. In fact, although it is difficult to make a quantitative assessment, HrpZ_{Pst} may actually be a slightly more potent elicitor of the HR than HrpZ_{Pst}.

However, $HrpZ_{Pst}$ appears to differ from the other HrpZ proteins in being deleterious to $E.\ coli$ cells when overexpressed and is possibly more unstable, making it difficult to purify large amounts of the protein. Since the glycine-rich region is the most obvious difference between $HrpZ_{Pst}$ and $HrpZ_{Pst}$ it is possible that it contributes to this phenomenon.

We were able to overcome this problem experimentally by using a tightly regulated T7 promoter/polymerase system, but never obtained quite the same level of expression we achieved with HrpZ_{Pss} and HrpZ_{Psg}. However, there remains the obvious question of how HrpZ toxicity is avoided by P. s. pv. tomato. One possibility would be that HrpZ is never expressed at levels high enough to affect the bacterium, even when it is induced in planta. Some indirect evidence for this hypothesis is provided by our examination of the DNA upstream of hrpZ_{Pst}. The ORF has a weak ribosome binding site, and we also observed that expression of cloned hrpZ from the lac promoter appears to be attenuated by the presence of cis-acting upstream sequences. A 9-bp GC-rich repeat upstream of hrpZ may be significant in this regard. Preliminary data from northern blotting experiments also indicate that premature transcription termination may take place when hrpA-hrpZ clones are expressed in E. coli (G. Preston, unpublished). A second possibility is that the location of the hrpZ gene in an operon with secretion genes ensures tight coupling of synthesis and secretion. Genes encoding extracellular proteins and secretion pathway components are often coregulated, but with a few exceptions involving the type I pathway, they do not lie within the same operon (Fath and Kolter 1993). A third possibility is that P. s. pv. tomato is more tolerant of high levels of HrpZ than is E. coli, or it possesses a means of keeping HrpZ in a nontoxic form while it is in the cell.

Further comparison with the Yersinia virulence system presents an intriguing possibility in this regard. It has been shown that secretion of certain "Yops" (the Yersinia pathogenicity determinants), involves chaperone proteins, small hydrophilic proteins which help keep the Yop protein in a translocation competent form and help target it for secretion (Wattiau et al. 1994). The genes encoding each chaperone are located adjacent to the gene encoding the corresponding Yop. Given the presence of several small ORFs of undetermined function in the pHIR11 hrp cluster, it is tempting to speculate that one of them, particularly hrpA, might encode a protein with chaperone function. There is a superficial resemblance between HrpA and Yersinia chaperones such as SycE. They are all small, hydrophilic, cytoplasmic proteins which lack a signal sequence, but there are no specific homologies. We are now constructing nonpolar mutations to test the role of HrpA in secretion. Preliminary results suggest that HrpA is not required for E. coli MC4100(pHIR11) to elicit an HR or secrete HrpZ (J. R. Alfano, unpublished), but in chaperone-mediated systems limited secretion of a protein will usually occur even in the absence of its chaperone, so it may be necessary to look quantitatively at secretion and accumulation of HrpZ to assess whether mutations in hrpA or other hrp genes have an effect.

The colinear relationship between several hrp and ysc genes.

From the sequence of the hrpZ operon it is clear that the parallels with the Yersinia type III secretion pathway extend beyond homologies of individual genes. The four genes downstream of hrpZ, hrpB-E, appear to be arranged colinearly with the region of the virC secretion operon from Yersinia that encodes YscI-L. The virC operon is a large operon containing 13 genes. yscA-yscM. several of which have been demonstrated to have a role in Yop secretion (Michiels et al. 1991). Of the four Yersinia genes with putative ho-

mologs in the hrpZ operon, only yscJ and yscL are known to have a role in secretion. An accompanying paper shows that five more hrp genes, downstream of the hrpH operon, are colinear with the yscQ-U genes in the virB operon of Yersinia (Huang et al. 1995).

It appears that a significant proportion of the type III secretion pathway described in *Yersinia* can be identified in *P. syringae*, and it seems likely that increasing parallels between the two systems will be found. In both systems the secreted proteins are involved with early events in the interaction with the host, and expression of secretion genes and virulence proteins is tightly coregulated. The secretion pathway seems to function in a similar way, as in both cases secreted proteins lack an N-terminal signal peptide and are not posttranslationally processed.

HrpZ and host specificity.

The function of HrpZ in compatible interactions is unclear. A likely role is the release of nutrients to the apoplast. Atkinson and Baker (1987a, 1987b) have proposed that the alkalinization of the apoplast caused by Hrp* bacteria (which occurs at a slower rate in compatible interactions) results in the leakage of sucrose and other nutrients to support bacterial growth. One of the key unanswered questions regarding the Psyringae HrpZ proteins is their role in host specificity. Compatible interactions leading to disease are distinguished by the absence of the HR. Host-differential elicitor activity would be one way to reconcile the production of HR-eliciting proteins by P. syringae and the phenomenon of host-specific compatibility. The failure of the PopA1 protein to elicit the HR in tomato, a host of P. solanacearum GMI1000, supports this concept (Arlat et al. 1994). Similarly, the isolated P. s. pv. syringae 61 HrpZ protein fails to elicit the HR in bean, although the significance of this is diminished by the fact that bean leaves appear insensitive to any harpins (He et al. 1993). To further explore this question, we infiltrated all three HrpZ proteins into the leaves of the host plants for each of the pathovars. The host plants of P. s. pv. syringae 61, and P. s. pv. glycinea, bean and soybean, respectively, are uniformly unreactive to HrpZ from both compatible and incompatible pathogens; however, tomato leaves proved to be highly sensitive to all three HrpZ proteins. Thus, our data argue against the hypothesis that host-differential activity of HrpZ proteins controls the host specificity of P. syringae pathovars.

If isolated $HrpZ_{Ps}$ elicits the HR in tomato, why does P s. pv. tomato not elicit the HR during pathogenesis? One possibility is that the response of tomato to HrpZ_{Psi} is qualitatively different than the response to HrpZ_{Fss} and HrpZ_{Fsf} despite manifestation of the same gross morphology. That is, the necrosis elicited by HrpZ_{Psi} is fundamentally different than the HR and does not involve associated defenses that stop the pathogen. We are now testing this possibility with probes for HR-specific transcripts. A second possibility is that HrpZ_{Ps:} production is regulated in a host-specific manner. However, hrpZ is clearly part of the Hrp regulon: hrpZ expression is transcriptionally linked with genes encoding components of the secretion pathway, the hrpZ operons in all three of these P. syringae pathovars have virtually the same hrp/avr promoter sequence, and expression of the hrpZ operon is likely required for pathogenicity. The conserved promoter sequences suggests that the hrpZ operon is regulated in P. s. pv. glycinea and P. s. pv. tomato by the same nutritional conditions and HrpR, HrpS, HrpL regulatory cascade described for P. s. pv. syringae and P. s. pv. phaseolicola (Grimm and Panopoulos 1989; Rahme et al. 1992; Xiao et al. 1992; Xiao et al. 1994; Xiao and Hutcheson 1994; Grimm et al. 1995). Whether differential expression of the Hrp regulon controls host specificity awaits determination. A third possibility is that the P. syringae pathovars produce host-specific suppressors of defense responses. This is supported by the observation that compatible pathogens do not trigger defense responses in host plants that are elicited by nonpathogens (Jakobek et al. 1993).

It is important to note that our data do not eliminate the possibility that the three HrpZ proteins actually have differential activity in host plants when delivered by living bacteria and that the HR observed may be an abnormal response resulting from the presentation of a high concentration of HrpZ in an artificial manner. In that regard, it is interesting that legumes, which appear insensitive to isolated harpins, respond to Hrp recombinant E. coli cells that secrete the same proteins (He et al. 1993). Experiments in which the hrpZ genes of P. syringae pathovars are switched or altered in their patterns of deployment should test more definitively the role of HrpZ in determining host specificity.

In conclusion, we have characterized an operon containing two components of the Hrp* system of P. syringae—a block of secretion-related genes that are conserved in eukaryotic pathogens in the genera Pseudomonas, Xanthononas. Erwinia, Yersina, Shigella, and Salmonella and a gene encoding an elicitor that is unique to plant pathogens. The elicitors found in the P syringae pathovars are a subfamily of a larger class that appears to be characteristic of plant pathogens, and which we postulate to have a role in releasing nutrients for bacterial utilization. Our challenge now is to determine how the various components of the Hrp system have been adapted to serve plant parasitism in the face of plant defenses.

MATERIALS AND METHODS

Bacterial strains and plasmids.

Bacteria and plasmids used in this study are shown in Table 2. Pseudomonads were routinely grown in King's B broth (King et al. 1954) at 30°C, but for certain experiments the hypderepressing minimal medium of Huynh et al. (1989), adjusted to pH 5.5, was used. E. coli was grown in LM (Sambrook et al. 1989) or terrific broth (Tartof and Hobbs 1987). Plasmids were introduced into bacteria by transformation (Sambrook et al. 1989) or electroporation (Gene Pulser, Bio-Rad).

Plant materials.

The plants used in this study were tobacco (Nicotiana tabacum L. 'Xanthii'), tomato (Lycopersicon esculentum Mill. 'Moneymaker'), and soybean (Glycine max L. 'Harosoy'). Plants were grown in a greenhouse or growth chamber at 23° to 25°C with a photoperiod of 16 to 24 h. Infiltration of plant leaves with HrpZ preparations was performed with blunt syringes as described (Huang et al. 1988).

DNA analysis and sequencing.

All DNA manipulations, except where specified, followed standard protocols (Ausubel et al. 1987; Sambrook et al. 1989). The hrpZ region of pHIR11 was subcloned into

pBluescript II (Huang et al. 1995). Two PsrI fragments of 2.2 and 2.4 kb from pCPP2201 and pCPP2200, respectively, were subcloned into pBluescript II SK(-) in both orientations. A series of overlapping nested deletions covering both strands was generated for each of the subclones using Erase-a-Base (Promega, Madison, WI). The deletions were sequenced from double-stranded templates using Sequenase version 2.0 (U.S. Biochemicals, Cleveland, OH) and forward and reverse M13 primers. Sequencing was completed using specific primers synthesized by Integrated DNA Technologies (Coralville, IA). In addition, the 3.7 and 3.6 kb SacI-EcoRI fragments, which overlap the PsiI subclones from pCPP2201 and pCPP2200. were also subcloned into pBluescript II SK(-) and sequenced using the ABI 373A DNA sequencer at the Cornell Biotechnology Program DNA sequencing facility and specific primers synthesized by IDT. Nucleotide and derived amino acid sequences were analyzed with the Genetics Computer Group Sequence Analysis Software Package (Devereaux et al. 1984). Homology searches against major sequence databases were done with the BLAST program (Altschul et al. 1990).

PCR amplification of hrpZ from P. s. pv. glycinea and P. s. pv. tomato.

The hrpZ genes of P. s. pv. glycinea and P. s. pv. tomato were amplified by PCR from the plasmids pCPP2202 and

pCPP2203, respectively. Reactions were performed using the PCR Optimizer kit (Invitrogen, San Diego, CA) according to the manufacturer's instructions. Reactions were overlaid with mineral oil and incubated in a Hybaid Thermal Reactor (Hybaid, Teddington, U.K.) using these cycle parameters: 2 min at 94°C, followed by 30 cycles of 1 min at 94°C, 2 min at 55°C. 3 min at 72°C, followed by a final incubation of 7 min at 72°C. The primers used for hrpZPsg were 5'and 5'-TACGGGATCCTTTGAGGAGGTTGTGATG-3' TACGCTGCAGTATC AGTCAGGCAGCAGC-3', and those for hrpZ_{Pst} were 5'-TACGGGATCCATGCAAGCACTTA ACAGC-3' and 5'-GGAACTGCAGCAAGCTCCGGCGA-TACAC-3'. All primers were synthesized by Integrated DNA Technologies, Inc. (Coralville, IA), and were designed to introduce a BamHI and a PsrI site at the 5' and 3' ends, respectively, of each amplified fragment.

The $hrpZ_{Psg}$ fragment from pCPP2202 was successfully amplified in all reaction buffers tested. The $hrpZ_{Psg}$ fragment from pCPP2203 was successfully amplified using reaction buffer B (reaction concentration 60 mM Tris-HCl, 15 mM (NH₂)₂SO₄, 2 mM MgCl₂, pH 8.5). PCR products of the expected sizes of 1.0 and 1.2 kb were purified from an agarose gel. digested with Psg and BamHI, cloned into pBluescript II, and then transformed into E. coli DH5 α F'lacI, yielding plasmid pCPP2255 carrying $hrpZ_{Psg}$ Plasmids containing

Table 2. Bacterial strains and plasmids used in this study

Designation	Relevant characteristics ^a	Reference or source
Escherichia coli DH5α	supE44 ΔlacU169 (080lacZΔM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1 NaF	Hanahan 1983; Life Technologies, Inc. Grand Island, NY
DH5c Flacf	F* proAB* lac[9ZAM15 zzf::Tn5[Km]/o80d lacZAM15 AdacZYA-argF)U169 endA1 recA1 hsdR17 (r, m,*) deoR thi-1 sapE442; gyrA56 relA1	Life Technologies Inc.
DI 21-DE21	F ompT hsdB _B (r _B m _B) dem ga! DE3	Novagen
BL21(DE3)	1 Original Roberts Co.	
Pseudomonas syringae	Wild type	Baker e: al. 1987
pv. syringae 61	Wild type	C. J. Baker
pv. glycinea race 4	Wild type, Rpf	D. E. Cuppels
pv. temato DC3000	with type, my	
Plasmids	Cloning vector, Amp	Stratagene
pBluescript II SK(-)	pUC19 derivative. Amp ^r	Schweizer 1991
pUCP19	T7 transcription vector. Amp'	Novagen
pET21(+)	T7 transcription vector, Amp	Tabor and Richardson 1988
pT7-6		New England Biolabs
LITMUS 28	Cloning vector, Amp ^r 25-kb cosmid containing P.s. pv. syringae 61 hrp cluster	Huang et al. 1988
pHIR11	25-kb cosmic contaming F.3. pv. syringer of mp closed	He et al. 1993
pSYH10	hrpZ _{Pst} ORF in pBluescript II 0.8-kb Pstl-Agel subclone from pHIR11, containing hrpB, in LITMUS 28	This study
pCPP2303	0.8-kb Psti-Agel subcione from prink it, containing http://	This study
pCPP2305	1.3-kb Sall-Sacl subclone from pHIR11, containing httpD, in pT7-6	This study
pCPP2200	pUCP19 carrying 10-kb partial Sau3A1 fragment of P. s. pv. glycinea DNA with httpZeij	This study
pCPP2202	2.4-kb Psrl subclone of pCPP2200 in pBluescript II: hrpApie and hrpZpie in expressed	
-	orientation with respect to P_{loc}	This study
pCPP2204	As pCPP2202 but with $hrpZ_{Pij}$ in reversed orientation to P_{lac}	This study
pCPP2206	7 A.b. Peri hmAse and hmZere subclane from DCPP2202 in DC121(+)	This study
pCPP2208	3 6.4h Sacl-FooR1 hmZer subclone from pCPP 2200 in pBluescript in	This study
pCPP2210	1.85-kb Bg/II-Psti hrpZ _{Pst} subclone from pCPP2202 in pc 1.21(+)	This study
pCPP2255	DCR amplified hm7er ORF in pBluescript II	This study
pCPP2201	at CD10 carrying 10-kh fragment of P. s. DV. tomato UNA with http2pp	This study
pCPP2203	2.2-kb $PstI$ subclone of pCPP2201 in pBluescript II; $hrpA_{Pst}$ and $hrpZ_{Pst}$ in expressed one entation with respect to P_{tot}	•
pCPP2205	As pCPP2203 but with $hrpZ_{p,i}$ in reversed orientation to P_{luc}	This study
pCPP2207	2.2-bh.hm7. subclone from pCPP2203 in pET21(+)	This study
	3.7-kb Saci-EcoRI subclone from pCPP2201 containing httpBCDE _{Ptt} in pBluescript II	This study
pCPP2209	3.7-kb SacI-EcoRI subclone from pCPP2209 in LITMUS 28	This study
pCPP2304 pCPP2211	2.0-kb Bg/II-Pstl hrpZ _{Pst} subclone from pCPP2203 in pET21(+)	This study

Amp' = ampicillin resistance: Nal' = nalidixic acid resistance; Rp' = rifampicin resistance.

PCR-amplified $hrpZ_{Psi}$ were found to be unstable and appeared to promote cell lysis.

HrpZ purification and analysis.

HrpZ was purified from E. coli as previously described (He et al. 1993) with the following modifications. Cells were lysed in either 5 mM 2-(N-morpholino) ethanesulfonic acid (MES), pH 5.5, or cell lysis buffer (50 mM Tris-HCl. 1 mM EDTA, 100 mM NaCl, pH 8.0). For some experiments the supernatant from heat-treated lysate was partially purified after sonication by ammonium sulphate precipitation (25 to 45% saturation), with desalting and concentration being performed with Centricon-10 tubes (Amicon). For experiments requiring highly purified HrpZ expressed in E. coli BL21(DE3), the supernatant was further purified by binding to phenyl-sepharose (Sigma) in the presence of ammonium sulphate (>30% saturation) and elution with 5 mM MES, pH 5.5, followed by electrophoresis through a native 15% polyacrylamide gel. The purified protein was then eluted from excised gel slices using an Elutrap apparatus (Schleicher & Schuell) or from crushed gel slices using a Micropure separator (Amicon). Protein concentrations were determined using Bio-Rad protein assay solution. HrpZ was also purified from heat-treated supernatants of P syringae grown in hrpinducing medium (Huynh et al. 1989) by ammonium sulphate precipitation (25 to 45% saturation) and desalting/concentration using Centricon-10 tubes. For infiltration into plant tissue. HrpZ preparations were diluted to various degrees with 5mM MES, pH 5.5. The amino-terminal sequence analyses were performed at the Cornell Biotechnology Program Protein Analysis Facility (HrpZ_{Pst}) and the University of Kentucky Macromolecule Structure Analysis Facility (HrpZ_{Fs}).

T7 expression and labeling of proteins in E. coli.

Proteins encoded by the *lirpZ* operon were expressed in *E. coli* BL21(DE3) by using the pET21(+) T7 expression system (Novagen). Conditions for isopropyl-β-D-thiogalactopyranoside (IPTG) induction of T7 RNA polymerase-dependent expression and labeling with L-[25] methionine were as described by Studier et al. (1990). After being labeled, cells were collected by centrifugation and then resuspended and lysed in SDS-loading buffer and the proteins resolved on an SDS-polyacrylamide gel. Gels were stained, dried and exposed to Kodak λ-ray film.

Nucleotide sequence accession numbers.

The nucleotide sequences reponded in this paper have been deposited in GenBank under accession numbers L41861 (P. syringae pv. tomato hrpA. hrpZ. hrpB. hrpC, hrpD. hrpE). L41862 (P. syringae pv. glycinea hrpA, hrpZ, hrpB). L41863 (P. syringae pv. syringae hrpA), and L41864 (P. syringae pv. syringae hrpB).

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hrp Genes of Phytopathogenic Bacteria

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1 Introduction

and Lindow in this volume). Among the 1600 different species known in the bacterial kingdom only a small number (about 80) are plant pathogenic and in most cases highly specialized with respect to the plant that can be attacked. Only a few of these species are gram-positive, e.g., Clevibacter ssp. and Straptomyces In nature plants are resistant to the majority of pathogens, and many bacteria live in close contact with the plant without causing any harm (see chapter by Beathe ssp. In this review I focus on subspecies of the gram-negative genera Erwinia, Pseudomonas, and Xanthomonas, which comprise the major bacterial plant

To be a successful pathogen the invading bacterium has to overcome the plant's defense. During evolution plant pathogenic bacteria have acquired multipie functions that enable them to colonize and multiply in living plant tissue. In nature, bacteria enter the plant through natural openings (stornata, hydathodes) or

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ment and severity of infection to different extents (see chapters by Dow and ing on the pathogen, i.e., these functions contribute to and modulate developpathogenicity. Obvious examples include degradative extracellular enzymes such as pectinases, cellulases, and proteases. When the corresponding genes are muteted, bacterial ability to invade plant tissues is more or less affected dependwounds. The bacterial armory contains a number of weapons that contribute to

C. DUM.

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Daniels, and Collmer and Bauer in this volume).

their mutant phenotype. $ilde{h}p$ genes are not only essential for pathogenicity on a normally a host for the particular pathogen (so called non-host). The incompatible presection is often associated with the induction of a hypersensitive reaction (HR) In addition, phytopathogenic bacteria possess a large number of genes needed for basic pathogenicity. These genes have been operationally defined as hp (hypersensitive reaction and pathogenicity; Linosnen et al. 1986) based on plant, i.e., the ability to cause disease in a compatible interaction, but also for the incompatible interaction with resistant host varieties or with plants that are not

clearly distinguished from typical disease symptoms. It is important to note that saprophytic or nonpathogenic bacteria such as Escherichia coli or Pseudopathogen multiplication and spread and thus in prevention of disease development. Under natural infection conditions the HR is microscopically small and can be induced by just one bacterial cell. Only when bacteria are introduced into plant tissue at high cell densities in the laboratory (about 10° colony forming units or more/ml) is the HR macroscopically visible as confluent necrosis and can be monas fluorescens do not induce the HR and are unable to multiply in plant tissue. infection (Krewent 1982; Lindsay et al. 1993). The HR results in prevention of production of phenolics and antimicrobial agents, e.g., phytoalaxins, at the site of in plants the HR is a rapid defense response involving localized plant cell death, the plant. In contrast to the use of the torm hypersensitivity in the animal field,

2 Isolation of hrp Genes and General Features

housekeeping functions were eliminated. A third characteristic of all htp mutants grow in minimal medium. This way mutants affected in genes for basic the ability to cause disease in susceptible plants and to induce the HR in resistant host or non-host plants (often tobacco). The second criterion for the isolation of genes specific for the plant interaction was to ensure that the mutants would still wild-type strain were inoculated into the host plant and screaned for loss of both methyl-N'-nitro-A-nitrosoguanidine) or transposon mutagenesis of a pathogenic bacteria except Agrobacterium. There are excellent reviews that describe the early work or focus more on one particular pathogen (Wulus et al. 1991; Bouchen et al. 1992). The majority of hrp genes have been identified by complementation of loss-of-function mutants. Mutants obtained by random chemical (e.g., Nnp genes have been isolated from all major gram-negative plant pathogenic is that they are unable to grow in the plant.

cosmid clones from a genomic library of the wild-type strain resulted in isolation of a cluster of hip genes localized in a 20 kb DNA region. This was the first induced mutants of P.s. pv. phaseolicola that had lost both the ability to induce indication that both the ability to cause disease and to induce the HR are mediated monas syringae pv. phaseolicola. Lavoaren and coworkers (1986) isolated Tn5halo-blight disease on been and the HR in tobacco. Complementation with The $h oldsymbol{p}$ genes were originally described for the bean pathogen $^{Pseudo-}$ by common steps in a "pathway"

1991; Lasy and Been 1992). This conclusion is based on DNA hybridization experiments and, of course, does not exclude the presence of genes with pathogenic bacteria do not contain hrp-related DNA sequences (Stall and were originally isolated from diseased plants as opportunists together with Rhizobium ssp. there seem to be no hrp gene equivalents present (BoxAs et al. (e.g., Huang et al. 1988; Lindgaen et al. 1988; Fig. 1C). In addition, genes with DNA Bonas, unpublished results). Interestingly, nonpathogenic xanthomonads that MINSAVAGE 1990; BONAS et al. 1991). In Agrobacterium turnefaciens or in strains of Fig. (A), Erwinia amylovora (Stenberder and Beer 1988; Barny et al. 1990; Waltens et al. 1990; Bayer and Been 1991), and several other pathovers of P. syringae homology, and in some cases functional homology, have been isolated from other species, e.g., the so-called wts genes from E. stewartii (Copum et al. 1992; LAsy and Been 1992), and a region containing pathogenicity genes from X.c.~
m pv.Fig. 1B), the Xanthomonas campestris pathovers campestris and vitians (Antat et giycines that complement hrp mutants of X.c. pv. vesicatotia (Hwanc et al. 1992); Since then hrp gene clusters have been cloned from a number of different al. 1991), translucens (WANEY et al. 1991), and vesicatoria (Bonas et al. 1991; bacteria. Examples include *Pseudomonas solanacearum* (Boucutr et al. 1987) functional homology to hrp genes in these species.

periplasmic membrane-derived oligosaccharide synthesis in $\it E.~coli.$ The $\it hrpQ$ have been shown to be functionally homologous to the products of the $\it E.~coli$ and hrpT genes from P.s. pv. phaseolicola (Mitter et al. 1993) will be discussed phaseolicola (Fellax et al. 1991). Besides being nonpathogenic and unable to mdoGH operon (Lousens et al. 1993). The mdoGH genes are required for induce the HR in tobacco, P. syringae hrpM mutuants are also affected in mucus production. The hrpM locus encodes two putative proteins which are similar and addition, several smaller hrp loci have been described that are not linked to the canipestris palliovars campestris (Kamoun and Kado 1990; Kamoun et al. 1992) and oryzee (Kawiyaii et al. 1993), and the hrpM locus in P.s. pv. syringae (NiePoLo et al. 1985; Микногионтих et al. 1988]. htpM is functionally conserved in pathovar In all of the cases mentioned above, the hrp genes are organized in clusters of 22-40 kb, and I will restrict most of this chapter to these large hrp clusters. In large cluster present in the same bacterium. These include a region in P. solanacearum (Huann et al. 1990), the hrpX locus that is conserved in X. later in this chapter.

3 Structural Organization and Relatedness of hrp Clusters

Genetic studies using transposon-induced insertion mutants in the respective bacterial wild-type strains revealed that the htp clusters contain at least six to eight complementation groups (Fig. 1). Some htp gene clusters have clearly been shown to be localized in the chromosome, e.g., in P.s. pv. phaseolicola (Rahme et al. 1991) and in X.c. pv. vesicatoria (Bonas et al. 1991), whereas in P. solanacesrum, the htp cluster is on a megaplasmid (Bouchen et al. 1987).

et al. 1992; Lasv and Been 1992). In addition, cross-complementation within a LINDGREN et al. 1988; ARLAT et al. 1991; BONAS et al. 1991; LABY and BEEN 1992). Due to sequence data it is now becoming more and more apparent that several hrp genes are conserved in all major gram-negative plant pathogenic bacteria (see below). Whether there are hrp genes that are clearly pathovar-specific can only be answered when complete sequence information becomes available for several Furthermore, at least some regions of the hrp clusters appear to be conserved on syningse, and also to E. amylovora (Boucher et al. 1987; Artar et al. 1991; Gough ierent strains of the same pathovar, as well as between pathovars or strains within a species, and in some cases also between species..However, the degree of conservation varies. DNA homology is high within pathovars of a given the DNA level between P. solanacearum and pathovers of X. campestris, P. subspecies indicated a high degree of functional conservation of h p genes (e.g., Striking similarities have recently been found between the hrp genes of pathogehs belonging to different genera. The first indication of homologies came subspecies, e.g., in P. syringse (Lindaren et al. 1988; Huang et al. 1991) and in X. campostris (Bows et al. 1991). The latter studies were recently extended by PCR using primers based on $\it hp$ sequences from X.c. pv. $\it vesicatoria$ (Leite et al. 1994). rom Southern hybridization studies. DNA homology was observed among dif-

4 Function of hrp Genes in Xanthomonas campestris pv. vesicatoria and Other Plant Pathogenic Bacteria

DNA sequence analysis of the hrp genes has revealed some important clues to their possible biochemical functions. One of the first genes sequenced was a regulatory gene, hrp5, from P.s. pv. phaseolicola (Grimm and Panopouros 1989). This gene as well as hrp8, a regulatory gene from P. solanacearum (Genin et al. 1992), will be discussed below in the context of gene regulation.

Since htp genes are environmentally regulated (see below), it was believed Since htp genes are environmentally regulated (see below), it was believed for a while that they would be encoding "alternative" proteins required for adaptation of the bacterium to the plant as the preferred environment. The recently discovered sequence similarities between several putative Hrp proteins and known proteins from other bacteria, however, led to a very different hypothesis, namely, involvement of Hrp proteins in protein secretion. We have

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Fig. 1A-C. Genetic and translational organization of the htp gene cluster of different plant pathogenic becara. A Xamhomonas campesuris pv. vesicationis is processium; and C Pseudomonas synngae by to a sequence sumitative between ORFs in priferent clusters the boxes are filled with the same pattern. For references, see text

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BS 82 B4

gene, hrp. 1. The hrp. operon contains eight ORFs, called hrp. 1-hrp. 86, etc. A region of about 4 kb between $\hbar m {\it E}$ and $\hbar r p {\it F}$ does not seem to be involved in the interaction with the plant because insertions in this region do not lead to a change transcriptional organization is depicted in Fig.1A. The loci hrpA and hrpB are transcribed from right to left; the other four loci and transcribed from left to right (SCHULTE and BONAS 1992a). According to the locus (hrpA-hrpF) we have numbered the ORFs consecutively. The hrpA locus appears to contain just one hrp sbility we predict 21 hrpgenes in the 25 kb hrpcluster of X.c. pv. vesicatoria. Their quences from this and other bacteria are not yet published, I will summarize our results and refer to the other phytopathogenic bacteria as I go along. Based on genetic analyses and the open reading frames (ORFs) with a high coding probsequenced the entire hrp cluster of X.c. pv. vesicatoria. Since most hrp sein phenotype (Bows et al. 1991).

nucleotide and magnesium binding domains. It is more similar to protein traffic preparation). The HrpB6 protein is a putativa ATPase with highly conserved ATPases than to proton pump ATPases, and the lack of membraite spanning resembles signal peptidase II sequences which are typical of lipoproteins (Fenselau et al. 1992). Experiments using radioactively labeled pelmitate are underway to test whether HrpB3 is a lipoprotein. Recently, both HrpB3 and HipA1 were shown to be localized in the X.a. pv. vesicatoria menibrane fraction using polyclonal antibodies (S. Fenselau, C. Marie, and U. Bonas, manuscript in sequence, suggesting an inner membrane localization (Fenserau et al. 1992). Both two (HrpB3) transmembrane domains, suggesting that a part of these proteins might be targeted to the outer membrane. The signal sequence of HrpB3 in X.c. pv. vesicatoria. A number of putative Hrp proteins are most likely associated with or localized in the bacterial membrane. For example, the HrpC2 HrpA1 and HrpB3 contain an NH2-terminal signal sequence and one [HrpA1] or What are the characteristics of the Hrp proteins? It should be noted that, ixcept for three proteins, expression of the other 18 has yet to be demonstrated protein sequence contains eight transmembrane domains but lacks a signal

and hrpD2 genes. Complementation studies indicated that part of the hrp region including different plant pathogens. High DNA sequence identity (more than Searches of the database revealed sequence relatedness of more than half of 90%) was found to a 2.7 kb fragment cerrying pathogenicity genes from X.c. pv. glycinns (Hwww. et et. 1992). The authors predicted two ORFs, whereas in X.c. pv. vesicatoria, this region contains three ORFs corresponding to the hrpC3, hrpD1 The X.c. pv. vesicatoria Hrp proteins with putative proteins in other bacteria, is colinear in the two pathovars of Xanthornonas (unpublished). domains suggests a cytoplasmic tocation (Fernset nu et al. 1992).

InpB regulatory gene from P. solanacearum which is not present in the 25 kb Inp similarity to X.c. pv. vesicatoria proteins (Table 1; Fig.1). One exception is the region or in the flanking region of the X.c. pv. vesicatoria hrp cluster as determi-The deduced amino acid sequences of hrp genes published from P. solanacearum (Goush et al. 1992, 1993; Genin et al. 1992) show significant ned by DNA sequence analysis and hybridization studies (T. Horns and U. Bonas,

that http genes in X.c. pv. vesicatoria are more closely related to P. solanacearum found in other bacteria nomenclature might become confusing. However, as long as the genes have not been shown to be functionally homologous, we will that was reported earlier (see above) is also seen on the protein level. It appears than to *P. syringae* and t*o Erwinia.* As more and more homologous *hrp* genes are solanacearum (Gough et el. 1993), whereas the hrplgenes from E. amylovora (Wei similarity to htpC2 from X.c. pv. vesicatoria. P.s. pv. syringae also contains a hpB3 related gene, called hpY, and a hpD2 related gene, $hpW(\mathsf{H.-C.}$ Huang. personal communication). Thus, the high degree of DNA sequence conservation unpublished). Furthermore, several of the proteins mentioned are conserved in (Table 1). The HrpA1 protein from X.c. pv. vesicatoria is 48% and 29% identical to proteins from P. solanacearum (HrpA; Goucн et al. 1992) and P.s. pv. syringae (HrpH; Huwo et al. 1992), respectively. HrpC2 from X.c. pv. vesicatoria is even more conserved, being 66% idential to the corresponding HrpO protein of P. and Been 1994) and from P.s. pv. syringae (Huanc et al. 1993) both show 62%other species (Flg.1), however, the degree of sequence similarity varies greatly continue to use these names.

1992a), and hrpZ (He et al. 1993) (see below), and of hrpJ from P.s. pv. syringae (Huans et al. 1993) in the X.c. pv. vesicatoria hrp cluster (unpublished; see Fig. 1). ria some genes are absent in the hrp region of more distantly related species. For example, there are no known homologs of the harpin genes hrp/\text{V/VEI} et al. Besides genes that are conserved among the major phytopathogenic bacte-

from X.c. pv. vesicatorialand two putative NoI proteins of Phizobium frediithat are Similarities of 50%-60% were found recently between HrpA1 and HrpB3 encoded by a cultivar specificity region. NoIT and NoIW mutants have a wider host range in nodulation of soybean (Meinivanor et al. 1993). In addition, the authors mention that release of proteins is affected.

their direct role in transport has yet to be demonstrated, it is believed that the Ysc and Ler proteins mentioned in Table 1 are parts of a special transport apparatus for Yop secretion. Similarly, Shigella flexnerisecrates virulence factors, called Ipa pethway from that previously described for protein secretion. The genes involved in Yscj, the Yops accumulate in the cytoplasm (Micuri's et al. 1991). Although invasion plasmid antigens), that are distinct from Yops but share the general NH, terminal signal peptide, and are secreted by using an entirely different in secretion are clustered on a 70 kb virulence plasmid. In case of a mutation, e.g., putative Hrp proteins are related to proteins in animal pathogens such as Salmonella, Stugella, and Yersinia ssp. Since the first similarities found were to the Ysc, Vir, and Lcr proteins from Yersinia ssp, this group of organisms bacame Humin et al. 1992). In Yersinia, these proteins are essential for the secretion of Since they are described in detail in the chapter by G.R. Cornelis, I will mention only a few important features. The Yops are hydrophilic proteins that lack a typical Last but not least, Table 1 summarizes the significant sequence similarities a "role model" for plant pathologists (Fenselau et al. 1992; Goucii et al. 1992; virulence factors, called Yops (Yersinia outer protein; MicHELS et al. 1990, 1991). which have been found to proteins from animal bacterial pathogens. A number of

HrpD2	^{\$} FOqnH	нФСЗ,	'‱H	FroC1 ²	'EBqiH		¹1A⊕H	ble 1. Sequence semiarities of
		····				•		anthomonas campestns gv. vesicatoria
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			(62%) H:bl,				*HqriH (4252) ·	seudomonas
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(%27)	(%ZS)		(%0 <u>/</u>)				*302Y (55,96)	eareaq siniz19
Spazers			•,		الحالاة ¹³ (1369)	(#EL)		ersinia pseudotuberculosis
(%4.5)			*AixM (%23)	(%55)	**LixM (% S2)	* Theq2 (#28)	"GwM	nenxell ellegin
(%19) adedS			*Avni (%78)	%95) (%95)	,	^{ट.} -1802 (%0१) '-143	** Dvni (# S2)	murumingyi silanomisi
EEby			FIPA™	FB/B3e		(% 59)		
#dild (%89)			(%£9)	(%29)		자(사용) (유용)		silitaus sulfoes
⁶ 2qoM	≪8qoM	-			•	,,।∃-8 (%E⊊)		iloo airtoinertas3
(%\$9)	(%67)		••••		₩ ²			STOYOTORS BINIMIS
			⁶⁸ 29)					Envelora amylovara
					자아 (유13)	•	15/MON 15/MON	mbeni murdosiriA
			(%SS) ## ^Q ##					Caulobacter crescentus

Similarities between deduced amino acid sequences of Hrp proteins from X.c.pv.vesicatoria and other proteins include conservative amino acid exchanges. Number include conservative amino acid exchanges. Number indicates percent similarity.

Superintensity numbers indicate references as follows:

J. Fewsellau et al. 1993; 2. Bonas et al., unpublished. 3. Gough et al. 1993; 10, Plavo et al. 1993; 11, Fields et al. unpublished, accession at L23495; 12, Volume et al. 1993; 13, Volume et

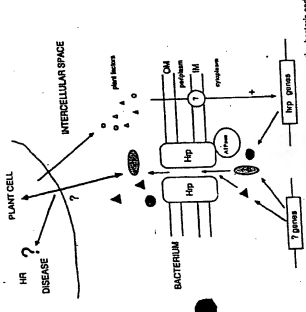


Fig. 2. Hypothotical model of call signaling botween gram-negative bacteria and plants indicating the proposed function of Hip proteins as an apparatus for protein secration. The model has been modified after FINELNU et al. 11932. Hip proteins may form a tunnel that enables the export of molecules such as virulence factors or swittence factors leading to either a hypersei sitive response IHRI or disease. Is these factors could be encoded by hip genes or genes unlinked to the large cluster. Both types of genes have been found to encode elicitors of the HRI (see text). The secretion of virulence proteins is hypithetical.

features mentioned above (Hale 1991; and see chapter by Parsn), this volume). Although S. typhimurium appears to possess a secretion system similar to that in Shigella, secreted invasion antigens have not yet been identified (Giuriavar and Octawav 1993; see chapter by Finit vi). As unpublished reports indicate that more and more genes in the animal pathogens are conserved, the data shown in Table and more genes in the animal pathogens are conserved, the data shown in Table 1 will soon be out of date. Proteins from other bacteria, e.g., E. coli, Bacillus, I will soon be out of bate. Proteins from other bacteria, e.g., E. coli, Bacillus, Caulobacter and from the mop region in E. carotovora (Muutoutwa) et al. 1993), Caulobacter and from the similar to Hrp proteins (Table 1). Most of these are have also been found to be similar to Hrp proteins (Table 1). Most of these are important for the assembly of the flagella, motility, or chemotaxis, again pointing, in my opinion, to a specialized secretion system rather than an involvement of hrp.

genes in chemotaxis.

These observations led us and others to propose a hrp-dependent secretion These observations led us and others to propose a hrp-dependent 1992; Van system in plant pathogenic bacteria (Fenseluu et al. 1992; Gouch et al. 1993). A modal is shown in Fig. 2 and raises certain questions, Guist (it we et al. 1993). A modal is being secreted by plant pathogenic bacteria? So e.g., if secretion occurs, what is being secreted by plant pathogenic bacteria? So e.g., a few proteins have been identified as elicitors of the HR but there is no lar, a few proteins have been identified as elicitors of the Southern in the evidence for secretion of virulence factors (see below).

5 hrp-dependent Secretion of Hypersensitive Response-Inducing Proteins

5.1 Harpin from Erwinia amylovora

An important feature of the isolated http clusters from both E. amylovora and P.s. pv. syringae is the ability of E. coli or Pseudomonas fluorescens transformants containing the cloned genes to induce the HR on tobacco (Huave et al. 1986; Been et al. 1991; see below). This has prompted to search for the HR-inducing activity within the respective gene clusters.

The first bacterial HR-inducing protein identified, designated harpin, is a cell anvelope-associated protein encoded by the hrpN gene of E. amylovora, a envelope-associated protein encoded by the hrpN gene of E. amylovora, a pathogen of pear and apple (Wet et al. 1992a). This harpin, is a glycine-rich and pathogen of pear and apple (Wet et al. 1992a). This harpin, is a glycine-rich and heat-stable protein that induces the HR in the non-host. Tobacco. The hrpN gene heat-stable protein that induces the normal host plant. Its function in pathogenicity, required for pathogenicity on the normal host plant. Its function in pathogenicity, required for pathogenicity on the normal host plant. Its function in pathogenicity, required for pathogenicity on the normal host plant. Its function in pathogenicity, any gene seems to be conserved among Erwinia sep. but that there is no DNA hrpN gene seems to be conserved among Erwinia sep. but that there is no DNA hrpN data described below suggest that the harping, protein might be secreted via the Hrp secretory apparatus, there is no published information available that demonstrates this.

5.2 Harpin from Pseudomonas syringae pv. syringae

E. amylovora, the function of harpines, in pathogenicity is unknown. Its product is secreted by P.s. pv. syringae in a HrpH-dependent way; HrpH is highly related to harpins harpin, and harpin, differ in their primary sequence, they have several features in common, e.g., a stretch of 22 amino acid that is similar in both proteins (He et al. 1993). Harpin, is also glycine-rich and heat-stable. As with harping of proteins involved in secretion in other plant and animal pathogens (Huang et al. occurs in natural infection is not clear. Interestingly, two short direct repeats in the COOH-terminus of harpings are essential for elicitor activity. Although the two containing an expression library, made using the cloned P.s. pv. syringae hrp cluster, were directly screened for HR-inducing activity on tobacco leaves. Two proteins were identified, one of which was an NH₃-terminal deletion of harpinps. , with even higher activity than the full size protein. Whether or not processing al. 1993; soe Fig. 1C and chapter by Collmer and Bauer). Lysates of E. coli clones which is encoded by the $\hbar pZ$ gene in the bean pathogen P.s. pv. syringse (HE et Using an elegant approach He and coworkers recently have identified harpinns. 1992; see Table 1).

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5.3 PopA from Pseudomonas solanacearum

An HR-inducing protein has been identified and characterized from *P. solana-cearum* culture supernatants, called Pop (Pseudomonas out protein; Arar et al. 1994). PopA1 and two shorter derivatives, PopA2 and PopA3, induce the HR in 1994). PopA1 and two shorter derivatives, PopA2 and PopA3, induce the HR in tobacco and in certain, but not all, *Petunia* lines. Like the harpins, the Pop proteins are also heat-stable and glycine-rich, however, the sequence is entirely proteins are also heat-stable and glycine-rich, however, the sequence is entirely proteins are also heat-stable and glycine-rich, however, the sequence is entirely proteins are also heat-stable and glycine-rich, however, the sequence is entirely and is dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent, I.e., the gene is part of the hip regulon. Mutations in popA do not dependent in Petunia is found it will be interesting (Fig. 1B). If a bacterial strain virulent towards Petunia is found it will be interesting (Fig. 1B). If a bacterial strain virulent towards Petunia is found it will be interesting (Fig. 1B). If a bacterial strain virulent towards Petunia is found it will be interesting (Fig. 1B). If a bacterial strain virulent towards Petunia is found it will be interesting (Fig. 1B). If a bacterial strain virulent towards Petunia is found it will be interesting (Fig. 1B).

the authors.

These exciting findings prove that certain Hrp proteins of P.s. pv. syringae
These exciting findings prove that certain HR elicitors (Fig. 2). They also
and P. solanacearum play a role in transport of HR elicitors (Fig. 2). They also
such a soland properties in the set of the set of

6 Regulation of Expression of hrp Genes

Expression of hrp genes is controlled by environmental conditions and has been studied on the RNA level as well as using transcriptional lusions to reporter genes studied on the RNA level as well as using transcriptional lusions to reporter genes such as the *E. coligenes* encoding β-galactosidase or β-glucuronidase. In general, expression of hrp loci is not detectable when the bacteria are grown in complex culture media. However, after growth of the bacteria in the plant, hrp genes are expressed. Attempts to mimic the conditions that the different bacterial species encounter in the plant tissue resulted in the finding that growth in minimal media without any plant-derived factor was sufficient to induce hrp genes. This has led to the speculation that the bacteria have to experience some kind of starvation to the speculation that the bacteria have to experience some kind of starvation conditions for full expression of hrpgenes. One of the first indications for hrpgene expression in vitro, and clearly a breakthrough, was a report on the hrp-dependent expression of an avirulence gene from the soybean pathogen P.s. pv. glycinea

(Hurwit et al. 1989).

Since the composition of minimal needla differs depending on the bacterium Since the composition of mindings will be summerized for representative studied, the most important findings will be summerized for representative pathogens. Parameters like carbon source, concentration of organic nitrogen and pathogens. Parameters like carbon been found to be important. High conphosphate, osmolarity, and pH have been found to be important.

centration of organic nitrogen generally appears to suppress hrp gene activation. Only two regulatory genes have been studied so far (see below). Interestingly, they both belong to different families of regulatory proteins.

6.1 Pseudomonas syringae

syringae occurs in the same medium as described by Huvnн et al. (1989); [Нилло et al. 1991; Хіло et al. 1992]. The authors showed hrp gene induction in the nonhrpL and hrpRS are only expressed to a very low level in M9 minimal medium and host plant, tobacco, but no data for the host plant. The P.s. pv. phaseolicola loci are induced at least 1000-fold when the bacteria are inoculated into the plant. This led to the conclusion that, at least for expression of hrpL and hrpRS, specific from P.s. pv. phasaolicola and was suppressed by TCA cycle intermediates such reaches the levels obtained in the plant (Ranme et al. 1992). A similar observation was made earlier for the avirulence gene avrB in P.s. pv. glycinea. Induction Expression of avrB is dependent on hrp genes homologous to hrpRS and hrpL. as citrate and succinate (HuvnH et al. 1989). hrp gene expression in P.s. pv. Expression of all seven hrp loci in the large cluster of P.s. pv. phaseolicola is suppressed in complex medium but induced in the plant. Induction occurs in the susceptible host plant as well as in the non-host, tobacco, suggesting that there is no plant species-specific molecule involved in control of host range (RAHME et el. 1992). Five complementation groups, hrpAB, hrpC, hrpD, hrpE and hrpF, can however, induction is affected by pH, osmolarity, and carbon source, and never occurred in a minimal medium containing fructose, mannitol, or sucrose. also be induced in M9 minimal medium containing sucrose as a carbon source, olant factors might be necessary (Ядние et al. 1992).

6.2 Regulatory Genes hrpRS and rpoN of Pseudomonas syringae pv. phaseolicola

The results on environmental factors inducing or suppressing hrp gene expression suggested that specific regulatory genes are involved in the control of hrp promoter activities. At least two loci are involved in positive regulation of the other promoter activities. At least two loci are involved in positive regulation of the other hrp loci of P.s. pv. phaseolicola hrp cluster (Fetrax et al. 1991). While there is no information published for hrpl., hrpRS has been sequenced. It contains two genes whose predicted protein products are 60% identical to each other (GRIMM and PANDOLOLOLOS 1989; Miller et al. 1993). The HrpS protein is similar to members of the NtrC family of regulatory proteins in gram-negative bacteria. Most NtrC-like regulatory proteins are members of two-component systems, with a sensor protein that in turn activates a response element by phosphorylation of a site in the conserved NH₂-terminal domain (Alenkent et al. 1989). The putative sensor component operating in hrp gene regulation has not been identified. It is postulated that HrpS is the activating protein, however, direct biochemical data

have not been presented. The tack of a typical NH₂-terminal domain in HrpS could indicate that a different mechanism may be involved in HrpS activation. Apparently, hrpSrelated sequences are also present in other bacteria, e.g., in P.s. pv. syringae (Hru and Hurcheson 1993) and in Erwinia amylovora (Been et al. 1993). E. stewartii contains a transcriptional regulator, WitsA, with 52% identity to HrpS of P.s. pv. phaseolicofa. The hrpS clone, however, was unable to functionally complement a wisA mutant (Frenenck et al. 1993).

complement a viscilination of the httpRS locus and the finding of -24/-12 consensus. The structure of the httpRS locus and the finding of -24/-12 consensus sequences upstream of httpRS indicated a possible role in transcriptional activation for transcription factor sigms 54, encoded by rooN(Gnium and Paviorou.os vation for transcription factor sigms 54, encoded by rooN(Gnium and Paviorou.os vation for transcription factor sigms 54, encoded by rooN(Gnium and Paviorou.os vation for transcription factor factor for the gene expression in P.s. pv. phaseolicola is a glutamine auxotroph and nonpathogenic. Whether of P.s. pv. phaseolicola is a glutamine auxotroph and nonpathogenic. Whether of P.s. pv. vesicatoria, rooN is clearly not involved in http gene expression X.c. pv. vesicatoria, rooN is clearly not involved in http gene expression and pathogenicity (T. Horns and U. Bonas, manuscript in preparation).

and pathogenicity (T. Horns and O. Bories), indirections of two new loci. Recently, Mitter et al. (1993) have reported the identification of two new loci. InpO and InpT, Irom P. s. pv. phaseolicola that affect activation of InpRS in trans. InpO and InpT, Irom P. s. pv. phaseolicola that affect activation of InpRS in trans. However, since InpRS is strongly induced in plants while both InpO and InpT are constitutively expressed, there must be more factors involved in Inp gene constitutively expressed, there must be more factors involved in Inp gene regulation. Strains carrying mutations in either InpO or InpT are amino acid regulation. Strains carrying mutations in either InpO and InpTare probably involved in auxothrophs (Imethiorine and tryptophan biosynthesis, respectively (Mitter et al. 1993). As methionine and tryptophan biosynthesis, respectively (Mitter et al. 1993). As stated above, such mutants would normally have been eliminated from the Inp stated above, such mutants would normally have been eliminated from the Inp

6.3 Conserved Sequence Boxes in Pseudomonas syringae

A conserved sequence, the so-called harp box (TG(A/C)AANC, Fritav et al. 1991). upstream of four hrp loci in P. s. pv. phaseolicola, was suggested to be involved upstream of four hrp loci in P. s. pv. phaseolicola, was suggested to be involved no positive regulation of expression. Similar motifs were described for the promoter regions of several P. syringae avirulence genes, the expression of which is dependent on hrpRS and on rpoN {HuviH et al. 1989; Salmenon and which is dependent on hrpRS and on rpoN {HuviH et al. 1989; Salmenon and vhich is dependent on hrpRS and on rpoN {HuviH et al. 1989; Salmenon and vhich is dependent on hrpRS and on rpoN {HuviH et al. 1989; Salmenon and vhich is dependent on hrpRS and on rpoN {HuviH et al. 1989}. These studies led to a Staskawiz 1993; Ihnes et al. 1993; Shen and Keen 1993). A harp box-related motif was also found upstream inducible {Shen and Keen 1993}. A harp box-related motif was also found upstream of transcription unit 3 in P. solanacearum {Goudh et al. 1993}.

There is no harp box sequence in *Xanthornonas hrp* gene promoters. Another There is no harp box sequence in the promoter region of *hrp* loci in *X*. c. pv. sequence motif that occurs in the promoter region of *hrp* loci in *X*. c. pv. sequence motify identified. This "PIP" (plant-inducible promoter) box *vesicatoria* was recently identified. This "PIP" (plant-inducible promoter) box has the sequence TTCGC-N15-TTCGC and occurs upstream of the "35 has the sequence in four out of six *hrp* promoters (S. Fenselau and U. Bonas, consensus sequence in four out of six *hrp* promoters (S. Fenselau and U. Bonas, binding motif.

6.4 Xanthomonas campestris

low concentration in phosphate. Both sucrose and methionine are needed for unpublished). A minimal medium was designed which would not suppress hrp gene induction. This medium, called XVM1, induces the hrpF locus (Fig. 1A) to high levels and differs from the other media described above, in particular by its efficient induction. It is not known whether a plant factor is necessary for activation of the other hrp loci, or if the XVM1 medium still lacks components or loci in X. c. pv. vesicatoria whereas the basal Murashige-Skoog culture medium did not. The inducing factor(s) could only partially be purified from TCM and was found to be smaller than 1000 dalton, heat-stable, organic, and hydrophilic within 1 h after transfer of the bacteria into TCM (S. Fenselau and U. Bonas, tomato cell suspension cultures (called TCM) induced expression of the six hrp (Schulte and Bows 1992a). De novo transcription of all hrp loci occurs rapidly as carbon source. No expression occurred in complex media or with high induced in the synthetic media tested so far. However, culture filtrates of sterile Expression of \emph{hm} genes in X. c. pv. $\emph{campestris}$ was determined after growth in vitro and found to be induced in a minimal medium with sucrose and/or fructose concentrations of organic nitrogen (Artat et al. 1991), in X. c. pv. vesicatoria, expression of the six hrp loci is induced in the plant but cannot be efficiently contains them in suppressing amounts (Schutte and Bonas 1992b).

6.5 Erwinia and Pseudomonas solanacearum

The hrp genes of Erwinia anylovora are rapidly induced in the non-host, tobacco, and more slowly in the host, pear. Several loci were induced in minimal medium with mannitol as a carbon source, Induction was suppressed by high concentrations of nitrogen and by glucose and was slightly temperature dependent (Welet al. 1992b).

In P. solanacearum, the hrp cluster was also induced in host and in non-host plants, as well as in minimal medium. The best carbon sources for induction of four of the six transcription units were pyruvate and glutamate while, as in other bacteria, casamino acids suppressed induction (Antar et al. 1992). The two rightmost hrp transcription units (5 and 6; Fig. 18) are constitutively expressed but can be induced under certain conditions (GENIN et al. 1992).

The only other gene reported to regulate htp gene expression is htpB from P. solanacearum. The gene is part of the htp cluster and appears to be a member of the AraC farnily of positive regulatory proteins. Interestingly, htpB is related to virF of Yersinia (Coinveius et al. 1989; Genin et al. 1992). The htpB gene positively regulates four of the six htp loci, as well as the popA locus, located outside of the htp cluster which encodes a protein secreted in a Hrp-dependent way (see above; An Ar et al. 1994). Whether the HrpB protein binds directly to htp promoters is not

yer kinowii. At this time one can only speculate whether the regulatory systems for *hrp* gene expression employed by *P. solanacearum* and *P. syringae* are really different

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mimic the dynamic nutritional situation that bacteria experience in their interaction with a plant for a short time. In mammalian bacterial pathogens, the expression of genes involved in virulence is also regulated in response to environmental cues rather than to specific host molecules. This subject has been reviewed recently (Mexalawos 1992 and in accompanying chapters), and I will only mention some important factors. In Yersinia, the vir and Icrgenes are regulated by reflect the situation in the plant. It is noteworthy that the in vitro culture will only plant factors as was described for the virulence genes of Agrobacterium (Winans 1992). Since the composition of the nutrients evailable to the pathogen in the plant is not known one can only speculate that the conditions described above gene expression in response to environmental cues. In conclusion, most hrp loci from different bacteria are inducible in a particular minimal medium. At this time or whether there is a global regulatory network thus allowing the line tuning of It cannot be ruled out that stimulation of *hrp* gene expression involves specific

not been described for any plant bacterium. In our laboratory no effect of calcium temperature (Conneus et el. 1989; see chapter by Conneus). A calcium effect has Shigella is affected by osmolarity and the later genes also by temperature (GALAN low calcium (low calcium response genes; Stratev et al. 1993) and by on htpF gene expression in XVM1 was observed (Schulte and U. Bonas, unpublished). Expression of invA of S. typhimurium of the mxi and ipa genes of and Cuntiss 1990; HALE 1991).

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The Enigmatic Avirulence Genes of Phytopathogenic Bacteria

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1 Action at the "Pathogenic Cusp"

armory of factors which determine successful colonization of that host. It is incumbent on each potential plant host, then, to evolve mechanisms to recognize some factor, preferably one produced at this pathogenic cusp, and to base resistance strategies on early recognition. Thus, an evolutionary tug-of-war is which not only the potential pathogen but also the host first sense and respond to each other. A successful plant defense response should be based on surveillance and interdiction before the pathogen has a chance to establish production of the genic growth strategies, "the pathogenic cusp" (Davia, 1994), is the point at The previous chapters have discussed how phytopathogenic bacteria can sense plant cell surfaces, and intracellular spaces. The switch from epiphyte to pathoand attendant function, as evidenced by induction of hip genes and subsequent production of various virulence and pathogenicity factors, some of which are hostspecific, some not. This reprogramming switch between epiphytic and pathoand respond to conditions present in a variety of microenvironments: soil, water, gen is apparently accompanied by fundamental reprogramming of gene activity

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MINIREVIEW

The Type III (Hrp) Secretion Pathway of Plant Pathogenic Bacteria: Trafficking Harpins, Avr Proteins, and Death

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INTRODUCTION

The ability of plant pathogenic bacteria to deliver deathtriggering proteins to the interior of plant cells was revealed in a rapid succession of papers in 1996 that transformed our concepts of bacterial plant pathogenicity. The breakthrough came with the convergence of work on Hrp systems and Avr proteins, an understanding of which requires an introduction to the most prevalent bacterial pathogens of plants, the cardinal importance of the Hrp pathway, and the paradoxical phenotype associated with avr genes.

Plant pathogenic bacteria in the genera Erwinia, Pseudomonas, Xanthomonas, and Ralstonia cause diverse, and sometimes devastating, diseases in many different plants, but they all share three characteristics: they colonize the intercellular spaces of plants, they are capable of killing plant cells, and they possess hrp genes. Many of these pathogens are host specific. In host plants, they produce various symptoms after several days of multiplication, whereas in nonhost plants, they trigger the hypersensitive response (HR), a rapid, defense-associated, programmed death of plant cells at the site of invasion (21, 43). With inoculum levels typically encountered in natural environments, the HR produces individual dead plant cells that are scattered within successfully defended healthy tissue (71). However, experimental infiltration of high inoculum levels (>106 bacterial cells/ml) results in macroscopically observable death of the entire infiltrated tissue, usually within 24 h (42). Pioneer screens for random transposon mutants with impaired plant interactions yielded a prevalent class that was designated Hrp, that is, deficient in both HR elicitation in nonhost plant species and pathogenicity (and parasitic growth) in host species (49, 56). This complete loss of pathogenic behavior results from mutation of any one of several hrp genes, which largely encode components of a type III protein secretion system (73). Because the capacity to elicit the HR is a convenient marker for the capacity to be pathogenic and these two abilities have a common genetic basis, the "simple" problem of HR elicitation is being studied as an entry to the larger problem of

A key part of the HR puzzle is that HR elicitation and the resulting limitation in host range can occur if the pathogen possesses any one of many possible aw (avirulence) genes that interact with corresponding R (resistance) genes in the host plant. Such "gene-for-gene" interactions result in recognition of the bacterium and the triggering of plant defenses. For example, Pseudomonas syringae pv. glycinea is one of over 40 P. syringae pathovars differing largely in host range among plant

species and is subdivided into races on the basis of their interactions with genetically distinct cultivars of its host, soybean. Those race-cultivar interactions involving matching bacterial av and plant R genes result in the HR and avirulence, i.e.; failure of the bacterium to produce disease. The R genes encode components of a parasite surveillance system and are crossed into crops from wild relatives by plant breeders for disease control. av genes are identified and cloned on the basis of the avirulence they confer on virulent races in appropriate test plants (39, 69). In most cases, it is not clear why plant pathogens carry av genes that betray them to host defenses but new insights into this question are discussed below.

Both hrp and avr genes were originally defined on the basis of the phenotypes they confer on bacteria interacting with plants. Molecular studies have revealed a functional relationship between the products of these two classes of genes and an underlying similarity with a key virulence system of several animal pathogens. Yersinia, Salmonella, and Shigella spp. transfer virulence effector proteins directly into animal cells via the type III pathway (16, 17, 62, 67, 84). Similarly, plant pathogens use the Hrp type III pathway to transfer Avr effector proteins to the interior of plant cells. The genetic dissection of type $\overline{\mathbf{III}}$ secretion systems is just beginning, and little is known of the mechanisms of protein translocation. In this review, we will describe (i) the recently completed inventory of genes directing type III secretion in plant pathogens and new insights into type III secretion mechanisms gained from research with Hrp systems, (ii) two classes of proteins (harpins and pilins) that are secreted by the Hrp type III pathway when plant pathogens are grown in media that mimic plant intercellular fluids, (iii) evidence that Avr proteins are delivered by the Hrp pathway directly to the interior of plant cells, and (iv) a resulting new paradigm for bacterial plant pathogenicity. The focus will be on quite recent work, and readers are referred to other reviews for a classic introduction to the HR phenomenon (43), earlier investigations of the Hrp system (11), avr genes (20, 46), and a wider perspective on bacterial virulence systems and plant responses (2).

Hrp PROTEIN SECRETION SYSTEM

hrp and hrc genes. hrp genes have been extensively characterized in four representative gram-negative plant pathogens: P. syringae pv. syringae (brown spot of bean), Erwinia amylovora (fire blight of apple and pear), Ralstonia (Pseudomonas) solanaceanum (bacterial wilt of tomato), and Xanthomonas campestris pv. vesicatoria (bacterial spot of pepper and tomato). Most of the known hrp genes in these strains are contained in chromosomal clusters of about 25 kb (Fig. 1). In at least some cases, the hrp clusters are sufficient to allow HR elicita-

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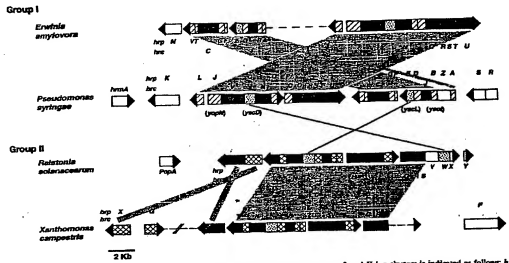


FIG. 1. htp gene clusters of four model plant pathogens. The distribution of each gene among group I and II htp clusters is indicated as follows: hrc genes, dark shading, htp genes that are conserved between groups I and II but show weaker similarity to Yersinia yet genes than hrc genes, stippling (the two lines between groups indicate homology); genes common to group I, diagonal lines; genes common to group II, hatching; genes for which no homologs have been reported, white. Dashed lines indicate gaps in the reported sequence of each htp cluster. The shaded bands between members of a group indicate colinear gene arrangements. Note that homologous htp genes have the same designation within group I but not within group II. Yersinia genes for which similarity has been noted with htp genes of R homologous htp genes have the same designation within group II but not within group II. Yersinia genes for which similarity has been noted with htp genes of R solanacearum (74), E. amplovora (10, 41), andor P. springe (36, 60) are in parentheses below the pair of group I htp clusters. The htp cluster of R solanacearum is solanacearum (74), E. amplovora (10, 41), andor P. springe (36, 60) are in parentheses below the pair of group I htp clusters. The htp cluster of R solanacearum of a megaplasming (12), but the others appear to be chromosomal. See reference 9 for previous designations of hrc genes and for references to all but the recent sequence reports in references 38, 41, and 81.

tion (but not disease) by nonpathogenic bacteria such as Escherichia coli and Pseudomonas fluorescens (8, 37).

Initial sequencing of the hrp clusters from R. solanacearum, X. campestris pv. vesicatoria, and P. syringae pv. syringae revealed homologies with components of the virulence protein (Yop) secretion system of Yersinia spp. (22, 29, 34), thereby suggesting the existence of a conserved "type III" protein secretion pathway in gram-negative pathogens of both plants and animals (65, 73). The near completion of these sequences has revealed further homologies and has led to two major changes in the nomenclature of hrp genes (9). First, those hrp genes that are broadly conserved in pathogenic Pseudomonas, Erwinia, Ralstonia, Xanthomonas, Yersinia, Salmonella, and Shigella spp. were redesignated hrc (HR and conserved) and given the last-letter designations of their Yersinia ysc homologs. The designations for Hrc homologs in various bacteria outside of the plant pathogen group are presented in Table 1. When referred to broadly, the term "hrp genes" is intended to encompass the hrc subset (9). Second, the hrp gene concept was widened to include homologous genes in plant pathogens where mutations do not lead to typical Hrp phenotypes. For example, mutations in hrp homologs result in loss of the Wts (watersoaking) phenotype in Erwinia stewartii (Stewart's wilt of corn) and reduced infectivity at low inoculum levels in Erwinja chrysanthemi (bacterial soft rot) (6, 23). Thus, the hrp genes appear to be universal among plant pathogenic Erwinia, Pseudomonas, Ralstonia, and Xanthomonas spp. and they control a variety of bacterium-plant interaction phenotypes in addition

Group I and II hrp clusters. The four hrp clusters that have been most characterized can be divided into two groups based on their possession of similar genes, operon structures, and regulatory systems (2). The hrp clusters of P. syringae and

E. amylovora are in group I, and those of R. solanacearum and X. campestris are in group II. In addition to the nine hrc genes, two hrp genes are conserved between the group I and II hrp clusters and show some similarities to yet genes (Fig. 1) (10, 36, 41, 74). It is likely that more of the present hrp genes will be discerned as belonging to the hrc category with additional data on the structure, function, and conservation of their products in both plant and animal pathogens. Nevertheless, some of the hrp genes appear to be completely different between the two groups, the arrangements of genes within some operons are characteristic of each group, and the regulatory systems are distinct (Fig. 1). A key difference in regulation is that group I hrp operons are activated by HrpL, a member of the ECF (extracytoplasmic function) subfamily of sigma factors (50, 78, 85), whereas most group II hrp operons are activated by a

TABLE 1. Hrc proteins of plant pathogenic bacteria and their animal pathogen and flagellar homologs

Plant pathogen	Yersinia	Salmonella	Shipella	Flagellar
protein	protein	protein	protein	protein(s)
HreC HreJ HreN HreQ HreR HreS HreT HreU HreV	YscC YscJ YscN YscQ YscR YscS YscT YscU LcrD	InvG PrgK SpaL SpaO SpaP SpaQ SpaR SpaR SpaS InvA	MxiD MxiJ Spa47 Spa33 Spa24 Spa9 Spa29 Spa40 MxiA	FIIF FIIL FIIN, -Y FIIP FIIQ FIIR FINB FINB

References for the sequences of hrc genes and all homologs are compiled in references 9, 25, and 74.

member of the AraC family, which is designated HrpB in R solanacearum and HrpX in X. campestris (27, 57, 82). However, hrp genes in both groups are generally repressed in complex media and expressed in plants and in media that mimic plant

intercellular fluids (11).

Functions of Hrp and Hrc proteins in type III protein se cretion. With the hrp clusters of four representative plant pathogens now almost completely sequenced, analysis of the functions of individual components is beginning. Nonpolar mutations have been constructed in most of the hrp and hrc genes in R. solanacearum and in some of the genes in P. syringae pv. syringae and E. amylovora (10, 15, 54, 77). The results suggest that the secretion apparatus requires all of the hrc genes (hrcQ awaits testing). The R solanacearum mutant analysis also reveals a requirement for hrpF, -W, -K, and -X (54). As discussed above, hrpF and hrpW have group I and possible ysc homologs. Thus, the Hrp type III secretion apparatus is likely composed of a core of 13 proteins, all but 2 of which appear to be broadly conserved. The predicted locations and functions of most of these proteins have been systematically presented for the R. solanacearum Hrp system (74), and they appear to be the same in X. campestris, E. amylovora, and P. syringae.

Sequence comparisons reveal that all of the Hrc proteins, other than HrcC, have a homolog involved in flagellum-specific export or early events in flagellum biogenesis (Table 1). The abilities of the presumably more ancient flagellar system to regulate the order (and possibly amount) of protein released and to secrete proteins in association with an extracellular appendage are properties that may be particularly important in the type III transfer of virulence proteins into host cells (18, 52). Plant pathogens offer several experimental advantages for exploring mechanisms of type III secretion and, indirectly, flagellum-specific secretion. The flagellum-specific and animal pathogen type III secretion systems have been difficult to study because many mutations pleiotropically disrupt production of the secretion apparatus and the secreted proteins. For example, the Yersinia pestis LcrD and Bacillus subtilis FlhA proteins (homologs of HrcV) were initially thought to have primary functions in regulation (14, 59). However, the unambiguous secretion phenotype of an E. amylovora hrcV mutant provided strong evidence that the primary function of members of this protein superfamily is in secretion (77). Plant pathogens offer other experimental advantages for exploring type III secretion mechanisms: defined subclones of ca. 25 kb are conveniently sufficient for Hrp-mediated secretion by E. coli and other model bacteria (31, 77), and hrc gene arrangements and mutant phenotypes suggest that translocation across the inner and outer membranes is partially separable in these bacteria (15).

In both group I and II hip clusters, the six hic genes predicted to encode a flagellum-derived system for Sec-independent translocation across the inner membrane (hrcN, -R, -S, -T, -U, and -V) are in operons other than that containing the one hre gene predicted to direct translocation across the outer membrane (hrcC) (Fig. 1 and 2). HrcC is a member of the PulD/pIV superfamily of outer membrane proteins, which are involved in type II protein secretion (PulD) and filamentous phage secretion (plV) (26). These proteins form homomultimers in the outer membrane which permit phage or protein exit and induce the psp (phage shock protein) operon (63). The HrcC protein of X campesurs pw. vesicatoria was the first member of the type III branch of this superfamily shown to induce the psp operon, thereby suggesting that the type III pathway also employs an outer membrane, channel-forming multimer (80). A P. syringae pv. syringae hrcC mutant accumulates some of the normally secreted HrpZ harpin (discussed below) in the periplasm, whereas a hrc U mutant accumulates

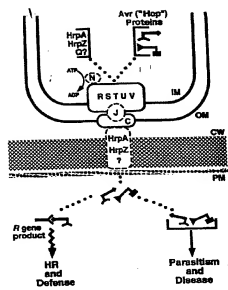


FIG. 2. Model for the delivery of parasite-promoting Avr proteins (i.e., Hop proteins according to a proposal discussed in the text) into plant cells by the Hrp type III secretion system (P. syringae example). To reach their targets, Avr proteins must cross the bacterial inner membrane (IM), outer membrane (OM), plant cell wall (CW), and plasma membrane (PM). Only Hrc components (flicated by their last letters in predicted subcellular locations) and proteins known to be secreted are shown. The location of hydrophilic HrcQ (HrcQ_A and HrcQ_B spr. syringae) is unknown, but the homologous SpaO is secreted by Salmonella spr. (25, 48). Four additional Hrp proteins, not shown, appear to be required for be secreted are shown. The location of hydrophilic HrcQ (HrcQ_A and HrcQ_B in P. syringae) is unknown, but the bomologous SpaO is secreted by Salmonella spp. (25, 48). Four additional Hrp proteins, not shown, appear to be required for secretion (see text). Dashed-line boxes indicate uncertainties about precise location. For example, it is not known whether HrpA or HrpZ penetrates the plant cell wall and whether these and/or other Hrp proteins trigger Aw transfer into plant cells by endocytosis. Secretion of HrpA and HrpZ is not dependent on plant cell contact, whereas secretion of Avr proteins apparently is. Once inside plant cells, multiple Avr proteins apparently promote parasitism collectively bunknown mechanisms (short arms denote weak phenotypes of virulence domains interacting with undefined host targets), unless any one of the proteins interacts with a host R gene product, thereby triggering the HR defense. Mustation of a host target, to diminish benefit to the parasite, and detection by the R gene surveillance system are likely evolutionary responses of plants to the bacterial deployment of a new virulence protein; coevolution would be expected to generate many aw and R genes in complex populations of plants and bacterial parasites.

the protein exclusively in the cytoplasm (15). Thus, the sequence-based prediction that separate inner and outer membrane translocator systems have been recruited to form the Hrp pathway is supported by a novel secretion phenotype revealing partial separation of these functions (15).

HARPINS, PILINS, AND OTHER PROTEINS SECRETED IN CULTURE BY THE Hrp SYSTEM

Harpins. Broadly defined, harpins are glycine-rich proteins that lack cysteine, are secreted in culture when the Hrp system is expressed, and possess heat-stable HR elicitor activity when they infiltrate the leaves of tobacco and several other plants. As is characteristic of proteins secreted by the type III pathway, harpins lack an N-terminal signal peptide. The first harpin was discovered in the culture fluids of E. coli cells carrying a highly expressed hrp cluster from E. amylovora (79). Because mutations in the harpin-encoding hrpN gene in E. amylovora strongly diminish HR elicitation in tobacco and pathogenicity in susceptible, immature pear fruits, harpin was initially thought to be the primary virulence protein traveling the Hrp pathway (79). Subsequent analysis of harpins from other bacteria has revealed that harpins differ substantially in their primary structure and their contribution to Hrp phenotypes, and their actual

function is unknown (4, 7, 19, 31).

The harpin genes of E. amylovora (hrpN) (79), E. chrysanthemi (htpNEch) (7), and R. solanacearum (popA) (4) are located adjacent to or near their respective hrp clusters, whereas the P. syringae hrpZ gene resides within a hrp operon (31). E. chrysanthemi hrpN mutants are reduced in infectivity at low inoculum levels and are unable to elicit the HR (7), but harpin gene mutations in E. amylovora CFBP1430 (a highly virulent strain) (5), R. solanacearum (4), and P. syringae (1) produce weak phenotypes or no phenotype. Thus, individual harpins do not appear to be necessary for elicitation of the HR by most bacteria. The potential role of harpins in determining host specificity is uncertain. PopA may be a host specificity factor because the isolated protein elicits the HR selectively in those plants in which R solanacearum also elicits the HR, whereas the isolated harpins from E. amylovora and three P. syringae pathovars trigger the HR in various plants in a manner that shows no relationship to bacterial host range (30, 31, 60, 79). Harpin activity may involve interactions with plant cell walls. The HrpZ harpin binds to the walls of intact plant cells but not to protoplasts, and it also fails to trigger HR-associated responses in protoplasts (33). The elicitor activity of harpins is unlikely to be enzymatically based because various fragments retain activity (1, 4, 45).

The function of the P. syringae HrpZ harpin is particularly

puzzling. Several observations suggest a simple, direct role for HrpZ in HR elicitation. HrpZ is the predominant protein secreted by the P. syringae Hrp system in culture (31, 88), the hrpZ gene is conserved in divergent P. syringae pathovars (60), and the isolated protein elicits an apparent programmed cell death in plants that is indistinguishable from the HR elicited by living bacteria (31). Furthermore, hpZ deletion mutations in the cosmid pHIR11 functional cluster of P. syringae pv. syringae hrp genes strongly reduce the HR elicitation activity of E coli cells carrying only pHIR11. The same mutation only slightly reduces the HR in P. syringae pv. syringae, but this can be explained by postulating the existence of a second harpin

encoded elsewhere in the bacterial genome (1).

However, other observations show that the relationship of HrpZ to HR elicitation is more complex. Mutation of hrmA (32, 35), which is in a variable region flanking the conserved hrp cluster in pHIR11, abolishes HR activity in tobacco without diminishing HrpZ synthesis or secretion (1). Thus, isolated HrpZ is sufficient to elicit an HR in tobacco Teaves but HrpZ produced by bacteria in plants is not. Instead, HrmA, with no apparent function in the Hrp secretion apparatus, is necessary for bacterial elicitation of the HR, and thus, HrmA appears to be the actual elicitor of the HR produced by bacteria carrying PHIRII. HrmA has several characteristics of an Avr protein (3). Avr proteins and the role of the Hrp system (and possibly harpins) in their delivery into plant cells will be discussed

HrpA pilin and other secreted proteins. P. syringae pv. tomato DC3000 secretes at least four proteins in addition to HrpZ into the medium in a Hrp-dependent manner (88). One of these is the 10-kDa product of hrpA, which forms a 6- to 8-nm-diameter "Hrp pilus" (61). A nonpolar hrpA mutant no longer elicits the HR in appropriate test plants, even when carrying an av gene known to interact with an R gene in the plant. It thus appears that the Hrp pilus is essential for the delivery of Avr signals (discussed further below). Although it is not known whether the Hrp pilus functions primarily in bacterial attachment or as a conduit for the delivery of bacterial

proteins across the plant cell wall, it is interesting that Agrobacterium tumefaciens requires a pilus similar in size (3.8-nm diameter) to transfer T-DNA and the VirE2 protein into plant cells (24).

H_{TP} DELIVERY OF AVR PROTEINS INTO PLANT CELLS

avr genes and their products. In fundamental contrast to the hrp genes, avr genes are scattered in their distribution among strains of plant pathogenic bacteria (20, 46). More than 30 bacterial av genes have been cloned from P. syringae and X. campestris, but until recently, characterization of the menagerie of encoded proteins has largely defined what these proteins do not do. Isolated Avr proteins do not elicit any responses when they infiltrate plant leaves. They do not appear to be secreted in culture and are hydrophilic proteins lacking N-terminal signal peptides or other recognizable secretion sign nals (properties consistent with potential secretion by the type III pathway). They do not have demonstrable enzymatic activ ity (with the exception of AvrD, which directs the synthesis of syringolide elicitors of an R gene-dependent HR [55]), and the majority of them do not contribute in an obvious way to parasitic fitness or virulence in the infection of cultivars lacking a matching R gene that would trigger the HR. However, there are several significant exceptions to the last point (20, 46) and there is growing evidence that Avr proteins have a primary function in virulence, even though the HR-triggering effects of Aw-R interactions are epistatic over these virulence functions. How Avr proteins might promote parasitism is mysterious, but support for such a primary role comes from observations that their action is dependent on the Hrp system and their site of action is within host cells. The next two sections address these points and provide evidence that the main function of the Hrp system is in the delivery of Avr-like proteins into plant cells. Hrp dependency of Avr phenotypes. avr genes have no phenotype when expressed in hrp mutant pathogens or in nonpathogenic bacteria like E. coli, which lack the Hrp system (highly expressed avrD is the sole exception to the latter point [40]). For many avr genes, especially those in P. syringue, one simple explanation is that their expression is dependent on Hrp regulatory factors (46). However, expression of avr genes from vector promoters does not obviate the requirement for a functional Hrp system. The recent finding that the functional cluster of P. syringae pv. syringae hrp genes carried on cosmid pHIR11 is sufficient to deliver heterologous avr gene signals indicates the fundamental interdependency of Hrp and Avr functions in bacterial elicitation of the HR (28, 58). A key property of pHIR11 enabling this discovery is that the cosmid confers on nonpathogenic E. coli and P. fluorescens the ability to elicit the HR in tobacco and several other plants, but it is ineffective in doing so in soybean and Arabidopsis. The simplest explanation is that hrmA, which is carried on pHIR11 and has several properties of avr genes (3), interacts with an unknown R gene in tobacco but with no R genes in soybean and Arabidopsis. This suggested that expression of appropriate avr genes in trans would enable nonpathogens carrying pHIR11 to elicit an R gene-dependent HR in soybean, Arabidopsis, and other plants. Indeed, this was observed with avrB (from P. syringae pv. glycinea) and five other P. syringue avr genes (28, 58).

The ability of pHIR11 to deliver avr gene signals requires HrcC (absolutely) and HrpZ (variably) (28, 58). The inability of HrpZ to support AvrB signal delivery when supplied exogenously indicates that the harpin has a role only when produced along with AvrB and therefore may be an extracellular accessory in the delivery of Avr proteins, as YopD is in the delivery of YopE (28, 62). Most importantly, these experiments reveal that a functional Hrp secretion system is required for the delivery of several av gene signals. Furthermore, the use of promoters different in strength and of epitope-tagged AvrB revealed that the requirement for a functional Hrp secretion system cannot be obviated by high levels of AvrB in the bacterial cytoplasm or by infiltration of leaves with purified AvrB at a level 1,000-fold higher than that required by living Hrp⁺ bacteria to elicit the HR (28). Thus, AvrB does not appear to act in the bacterial cytoplasm or in leaf intercellular spaces. These observations strongly support the hypothesis, depicted in Fig. 2, that the type III protein secretion system in plant pathogens, as in animal pathogens, is capable of deliver-

ing bacterial proteins into host cells.

Demonstrations of Avr action in host cells. Bacterial transfer of Avr proteins into plant cells has not been observed directly. However, there is evidence that several of these proteins are biologically active when produced within plant cells, that the HR-triggering activity of one of them is dependent on physical interaction with its cognate plant R gene product, and that the activity of another is dependent on localization to the plant cell nucleus. AvrB action in plant cells was demonstrated with Arabidopsis plants carrying the cognate RPM1 R gene (28). An Arabidopsis rpm1 mutant was transformed with avrB and crossed with a wild-type line, thus producing seedling progeny carrying both avrB and RPM1 which died soon after germinating. One symptomless rpm1 mutant transgenic plant was obtained; this individual expressed relatively low levels of an avrB construct carrying the PR-1a plant protein signal peptide, with the likely consequence that the plant cytoplasm would be exposed only transiently or to low levels of AvrB. The properties of this survivor suggest that plants are sensitive to AvrB even in the absence of a functional matching R gene and that vanishingly low levels of the protein are sufficient to elicit the HR in the presence of a complete R gene. A biolistic, transient expression assay revealed that avrB lacking a signal peptide (and therefore localized to the plant cytoplasm) was lethal to Arabidopsis leaf cells carrying RPM1 but not to those lacking the R gene (28). This approach was extended with avrRpt2 (from P. syringae pv. tomato) (47). Similarly, an A. tumefaciens transient expression system was used to deliver avrPto (from P. syringae pv. tomato) and avrBs3 (from X. campestris pv. vesicatoria) into plants, resulting in an R gene-dependent HR in all cases (66, 70, 72). Thus, whereas no bacterial Avr protein has been observed to have an effect when delivered exclusively to the surface of plant cells, all four of those tested elicit an

R gene-dependent response when expressed inside them.

The simplest model for the molecular basis of gene-for-gene HR elicitation predicts physical interaction between the protein products of cognate aw and R genes. This has been observed with the bacterial AvrPto and plant Pto proteins; mutations in the molecular partners that diminish physical interaction in the yeast two-hybrid system also diminish biological function (66, 70). Because AvrPto action requires a functional Hrp system in either P. syringae pv. tomato (64) or nonpathogens carrying the pHIR11 hrp cluster (28, 58) and it involves physical interaction with a cytoplasmic target in the host, the Hrp-mediated transfer of AvrPto into plant cells

seems certain.

While many bacterial Avr proteins appear to be targeted to the host plant cytoplasm, members of the AvrBs3 family in *Xanthomonas* spp. are targeted to the host nucleus. These proteins carry functional nuclear localization signals (NLS) in the C-terminal region (72, 86). When fusions of this C-terminal region and a *uidA* reporter are transiently expressed in onion epidermal cells by biolistic bombardment, β -glucuronidase ac-

tivity is localized to the nucleus (72, 86). Deletion of all three of the NLS sequences abolishes nuclear localization in the biolistics assay and HR elicitation by X campestris pv. vesicatoria cells in pepper plants carrying the Bs3 R gene, and both of these abilities can be restored by substitution of the simian virus 40 large-T antigen NLS (72). These results suggest that the Bs3 product must also be localized to the nucleus, but because this R gene has not been cloned, this awaits confirmation.

Gaps in our knowledge of the Hrp pathway and the inventory of its protein traffic. Although the rings of evidence that the Hrp system transfers Avr proteins into plant cells are collectively strong, there are formal gaps in each. (i) In the system explored in the most detail, AvrPto-Pto, physical interaction between the bacterial and plant proteins has not been demonstrated in vivo, and a second host protein, Prf, is required for AvrPto-Pto-mediated HR elicitation. Furthermore, all of the other cloned plant R genes that interact with known bacterial avr genes resemble Prf (a nucleotide-binding site leucine-rich repeat protein) rather than Pto (a kinase) (68). (ii) R proteins appear to be present at vanishingly low levels, and none has been directly observed in the cytoplasm, although RPS2 localizes to the cytoplasm-equivalent fraction in a rabbit reticulocyte dog pancreatic microsome in vitro translationtranslocation system (47). (iii) Similarly, Avr proteins appear to be effective at vanishingly low levels (28) and immunogold labeling and electron microscopy of infected plant tissues has revealed their presence only in bacterial cells (13, 87). (iv) Finally, no Avr protein has been directly shown to be translocated out of the bacterial cytoplasm in culture by the Hrp system. It is worth noting that the A. tumefaciens VirE2 protein has never been observed to be transferred into plant cells, although the indirect evidence for its action within plant cells seems irrefutable (89).

Many (if not most) of the genes encoding proteins that are transferred into plant cells by these bacterial pathogens probably await discovery. Systematic completion of the inventory is thwarted by two problems. First, the contribution of the genes to virulent interactions may be too subtle for detection in mutant screens, and cognate R geries that would reveal Avr phenotypes when the bacterial genes are heterologously expressed may be unknown or nonexistent. Second, no plant signals or regulatory mutants have been found that permit bacteria to secrete these proteins in culture, although harpins, pilins, and possibly other proteins that serve the type III secretion system are secreted in culture. A critical feature of the type III protein secretion system in Yersinia spp. is its capacity to withhold full secretion of virulence proteins until contact with the host cell (18). The fact that nonpathogens carrying the pHIR11 functional hrp cluster secrete HrpZ but not AvrB in culture (28) indicates that the genetic information for this expected regulatory step is carried within the hrp cluster and is therefore subject to discovery through systematic analysis of the hrp genes. Obtaining Avr protein secretion in culture is important because (i) it is likely to be associated with structures that normally are used to penetrate the plant cell wall (and possibly trigger host cell endocytosis) and therefore will yield clues to the transfer process and (ii) it will allow proteins targeted to the host to be systematically characterized through identification of novel proteins in the medium. The exploration of DNA sequences flanking hap clusters also should be useful in this search because of the growing evidence that these regions are enriched in genes whose products probably travel the Hrp pathway (51, 53, 54).

A new designation for effector proteins that are delivered by the Hrp system to plant cells would be useful: Avr appears to be inappropriate because some of the encoding genes may have no Avr phenotype and the primary function of Avr proteins is almost certainly in virulence, not avirulence. One proposal is to designate new members of this class Hop (Hrpdependent outer protein) and to add a four-letter suffix identifying the bacterial species, pathovar, and gene, based on the current system for uniform nomenclature of avirulence genes (3, 75). For example, the gene encoding a newly found P. syringae pv. syringae protein in this class would be designated hopPsyA. Hop is analogous to the Yop (Yersinia outer protein) designation for proteins secreted by the prototypical Yersinia type III secretion system but is broadened here for consistency with the use of Hrp and Avr for plant pathogens in all genera.

A NEW PARADIGM AND FUTURE EXPLORATIONS

Pathogenesis based on the Hrp delivery of Avr-like (Hop) proteins into host cells (depicted in Fig. 2) provides a simple and unifying explanation for many characteristics of plant pathogenic Erwinia, Pseudomonas, Xanthomonas, and Ralstonia spp. (2). These include the one-to-one relationship between bacterial cells and HR-responding plant cells (expected with contact-dependent secretion), the gene-for-gene interactions of pathogen races and host cultivars (expected if avr and R gene products can directly interact within host cells), and the enormous diversity in host range and other pathogenic attributes among closely related strains (expected with a pool of horizontally transferable and interchangeable genes whose products can either promote or betray parasites in coevolving hosts). The latter point is particularly relevant to P. syringae and X. campestris, which are divided into more than 40 and 140 pathovars, respectively. And it is consistent with the location of many avr genes on plasmids and the ability of avr genes to function with heterologous Hrp systems (20). In this regard, one potential difference between the type III systems of animal and plant pathogens is noteworthy. In animal pathogen type III systems, the secretion of many effector proteins requires customized chaperones, which are often encoded by genes linked to effector genes (76). The ability of many isolated avr genes to function heterologously in other pathogens or in nonpathogens carrying the pHIR11 functional hrp cluster suggests that Avr protein delivery does not require specific chaperones or that a promiscuous chaperone gene exists within the hrp cluster.

This new model of plant pathogenicity invites several fundamental questions in plant pathology and pathogenic microbiology in addition to those discussed above regarding the Hrp system and the identification of its traffic. How do Hrp-delivered proteins alter host metabolism to promote bacterial growth in plant intercellular spaces? How is host specificity determined at the pathovar-host species level? That is, are avr-R gene interactions important here also, as suggested by the discovery of novel avr genes through expression in heterologous pathovars (44, 83), or do Avr-like proteins have important positive effects in bacterial adaptation to host species? Given the use of homologous secretion systems, how similar are the functions of the virulence proteins that plant and animal pathogens transfer into their hosts? Sequence similarities involving secreted Yesinia proteins have been noted only between YopN and YopJ and the E. amylovora HrpJ and X. campestris pv. vesicatoria AvrRxv proteins, respectively (10, 46). Since YopN appears to be an extracellular component of the secretion system and the effector activity of YopJ is unknown, this key question remains unanswered. Further comparisons should give us a broader perspective on the evolution of bacterial pathogenicity and may lead to unanticipated controls for diseases of both plants and animals.

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Isolation of the hreX Gene Encoding the HR Elicitor Harpin (Xcp) from Xanthamonas campestris pv. pelargonii. S. SWANSON and Z-M. Wei. EDEN Bioscience Corporation, Bothell, WA 98011 USA. Phytopathology 90:S75. Publication no. P-2000-0537-AMA.

This study reports the isolation of a gene encoding a proteinaceous HR elicitor from Xanthomonas campestris pv. pelargonii, Xcp. The HR elicitor exhibits a high potency for eliciting HR in tobacco. Treatment of the Xcp HR Elicitor with proteases resulted in a loss of HR activity. Degenerate oligonucleotides were designed based on amino acid sequences obtained from the purified HR elicitor and used to screen a Xanthomonas campestris pv. pelargonii genomic library. An open reading frame, ORF, was identified consisting of 381 base pairs that encoded a protein of 126 amino acids. The ORF initiated with a typical methionine start codon and was preceded by a putative ribosome-binding site. The ORF was designated as the hreX gene, encoding the HR elicitor harpin (Xcp). HreX has a molecular weight of 13.3KD, a theoretical pI of 3.8 and is glycine rich. Further studies of harpin (Xcp) and its bioactivity are currently underway.

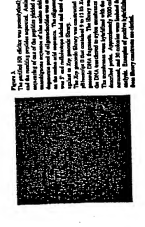
Isolation of the hreX Gene Encoding the HR Elicitor Harpin_{xp} from Xanthamonas campestris pv. pelargonii. S. SWANSON, Z-M. Wei.

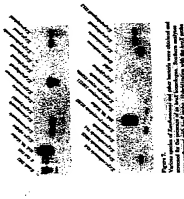
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GENE ISOLATION PROTEINACEOUS HR ELICITOR





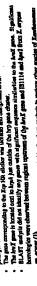








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Unified nomenclature for broadly conserved hrp genes of phytopathogenic bacteria.

Genes of plant-pathogenic bacteria controlling hypersensitive response (HH) elicitation and pathogenesis were designated 'hip' by Lindgren et at in 1986 (J Baoteriol 168: 512-522). hrp genes have been characterized. in several species of the four major genera of Gramnegative plant pathogens, Erwinia, Pseudomonas, Rajstonia (a new proposed genus including Pseudomanas: solanacearum) and Xanthomonas. To date, hrp genes have been found mainly in large clusters, and they have been shown to be conserved physically and, in many cases, functionally among different bacteria. Hybridization studies and genetic analyses have revealed the presence of functional hip genes even in species that are not typically observed to elicit an HR; such as Erwinia chrysanthemi and Erwinia stewarth, suggesting that hip genes may be common to all Gram-negative plant pathogens. possibly excluding Agrobacterium spp. Current knowledge of hirp genes has been reviewed by Bonas (1994, Curr Top Microbiol Immunol 192:79-98) and by Van Gijsegem et al. (1995, In Pathogenesis and Host-Parasite Specificity in Plant Diseases: Histopathological, Biochemical, Genetic. and Molecular Basis. Volume 1. (Kohmoto et al., eds); Oxford: Pergamon Press, pp. 273-292).

The nucleotide sequences of four hap gene clusters, those of Raistonia solanacearum (previously P. solanacearum) (Genin et al., 1992, Mol Microbiol 6: 3065-3076; Gough et al., 1992. Mol Plant-Microbe Interact 5: 384-369; Gough et al., 1993, Mol Gen Genet 239: 378-392; Van Gijsegem et al., 1995, Mol Microbiol 15: 1095-1114), Erwinia amylovora (Bogdanove et al., 1996, J Bacteriol 178: 1720-1730; Wel and Beer, 1993, J Bacteriol 175: 7958-7967; Wel and Beer, 1995, J Bacteriol 177; 6201-6210; Wel et al., 1992, Science 257: 85-88; S. V. Beer, unpublished). Pseudomonas syringae pv. syringae (Huang et al., 1992, J Bacteriol 174: 6878-6885; Huang et al., 1993, Moi Plant-Microbe Interact 6: 515-520; Huang et al., 1995, Moi Plant-Microbe Interact 8: 733-746; Lidell and Hutcheson, 1994, Mol Plant-Microbe Interact 7: 488-497; Preston et al., 1995, Moi Plant-Microbe Interact 8: 717-732; Xiao et al., 1994, J Bacteriol. 176: 1025-1036), and Xanthomonas campestris pv. vest. catoris (Fenselau et al., 1992, Mol Plant-Microbe Interact 5: 390-396; Fenselati and Bonas, 1995, Mol Plant-Microbe Interact 8: 845-854; U. Bonas, unpublished), have been largely determined. These clusters each contain-

more than twenty genes, many of which encode components of a novel protein-secretion pathway designated type III. It has been shown directly that various extracellular proteins involved in pathogenests and defence elicitation by plant-pathogenic bacteria utilize this pathway (Ariat et al., 1994, EMBO J 13: 543-553; He et al., 1993, Cell 73: 1255-1266; Wel and Beer, 1993, Ibid.), and the pathway is known to function in the export of virulence factors from the animal pathogens Salmonella lyphimurium. Shigella flexneri, and Yersinia entercollica, Yersinia pestis, and Yersinia pseudotuberculosia (for reviewa, see Salmond and Reeves, 1993, Trands Blochem Sci 18: 7-12; and Van Gijsegem et al., 1993, Trends Microbiol 1: 175-180). Nine type III secretion genes are conserved among all tour of the plant pathogens listed above and among the animal pathogens. Based on sequence analysis and some experimental evidence, they are believed to encode one outer-membrane protein, one outer-membrane-associated lipoprotein, five trans-membrane proteins, and two cytoplasmic proteins, one of which is a putative ATPase. All of the predicted gene products, except the outer-membrane protein, show significant similarity to components of the flagellar biogenesis complex (for reviews see Biair, 1995; Annu Rev Microbiol 49: 489-522; and Bischoff and Ordal, 1992, Mol Microbiol 6: 23-28). We herein refer to the hrp-encoded type III pathway as the Hrp pathway.

Because trip genes have been characterized independentity in diverse plant-pathogenic bacteria, hrp gene nomenclature differs in different species, and it is notalways consistent even within the same organism. Difterent designations are used for homologous genes, and, even worse, the same designation is used for different genes in different organisms. For example, hrpt of E. amylovors is homologous with hrpC2 of X campestris pv. vesicatoris and hrpO of R. solanacearum, and the homologue in P. syringae pv. syringae appears in the literature both as hrpl and as hrp.12. Also, thrpN' in Ft. solenacearum designates a secretion-pathway gens, whereas in E. amylovora, 'hrpN' designates the gene encoding the elicitor harpin. Furthermore, in many bacteria the number of known hip genes approaches 26. In anticipation of exhausting the alphabet, some authors chose to designate hip genes with a letter and a number, creating the potential for confusion of distinct genes with alleles of the same gene. For hrp gene researchers, the current nomenclature is at best inconvenient; for other scientists, it is bewildering.

Another problem exists: accumulation of knowledge about the structure of hip loci has outpaced the accumu-

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letion of information regarding the specific functions of individual genes. Typically, hrp loci have been identified by polar, transposon mutagenesis. Conceivably, a particular gena within an operon required for the Hrp phenotype may not be a strict Hrp determinant, but may play a more subtle role. Moreover, even phenotypes of mutations in well-characterized hip genes are not the same in all pathogens. For example, although the hrpN gene of E. amylovora is required for pathogenesis of pear truit, the homologous gene in E stewarti (D. L. Copilin, unpublished). is dispensible for pathogenicity of corn. In the macerogenic bacterium E chrysanthemi, even polar mutations that disrupt hip secretion altogether only reduce the apparent frequency of successful infection initiation (Bauer et al., 1994, Mol Plant-Microbe Interact 7: 573-581). Thus, the designation 'hrp' in its strict sense, i.e., meaning required for the HR and pathogenicity, is not uniformly applicable.

At the 7th International Congress on Molecular Plant-Microbe Interactions held in Edinburgh, Scotland in 1994, a committee of hip researchers and others was formed to address these problems. We, the committee members, agreed upon a system to standardize names for the subset of hip genes that are broadly conserved, and agreed to broaden the definition of the 'hrp' designation, as tollows.

For the subset of hrp genes that are broadly conserved. the new, unique, lower-case symbol 'hrc' will be used. The 'hr' of hip has been retained in order to evoke that name, and the 'c' has been added to denote 'conserved.' The upper-case designations will correspond to those of the type III secretion genes of Yersinia spp. (for a review, see Forsberg et al., 1994, Trends in Microbiol 2: 14-19), yscC, yscJ, yscN, yscQ-U, and lcrD, except that the lcrD homologues will be designated 'hrcV' to avoid confusion : of these as homologues of yscD, which is another, less, well-conserved type III gene of Yersinia spp. We request that Yersinia researchers omit the letter 'V' in naming any new yee genes that might be discovered. The yes

nomenciature was chosen as a standard for revising hip gene names for its convenient uniformity, and because, of all the genes that comprize the several known type III systems, the Yersinis genes show the highest degree of sequence similarity to the type III (hrp) genes of plant pathogens. The new names for the nine genes are given in Table 1, along with the current names in R. solanscearum, E. amylovora, P. syringes pv. syringes, and X. campestris pv. vesicatoria, and the names of homologues involved in flagellar biogenesia.

in designating genes as the, proadily conserved genes were defined as being present among the hip genes of at least one representative species of each of the four plantpathogenic genera discussed here and among the type ill genes of each of the enimal-pathogenic species S. typhimurium, S. flexneri, and the three yersinise. Gene tarnilles were defined based on pairwise sequence alignments. Any two genes were considered homologous if a sest-HT alignment (Devereux et al., 1984; Nucl Acids Res 12, 367-395) of the predicted amino acid sequences using detault. parameters yielded a quality econe at least five times the standard deviation above the mean quality score of 100 alignments, for each of which one of the sequences had been randomized prior to alignment (Doolittle, 1986, Of URFs and ORFs: a Primer on How to Analyse Derived Amino Acid. Sequences. Mili Valley, California: University Science Books).

Genes that did not meet the criterion for the 'hrc' designation will remain 'hip'. We have chosen to use this criterion until more data regarding structure and precise function of the products of the hip and other type III genes becomes available. Some of the genes that clid not meet the criterion In fact may be common to Relatonia, Erwinia, Pseudomonas, and Xanthomonas, and have homologues in the animal pathogens, yet may be sufficiently diverged to obscure obvious homology by direct sequence comparison. As structural and functional data accrue, such relationships may become clear, and the list of hir genes

Table 1. Current names and new, unified names for the broadly conserved hip games of R. solanscourum, E. amylovors, P. syringse pv. syringse. sicularia. Homologues that function in flagellar biogenesis are given in the bottom row.

Table 1. Current name:	vesicaloria. Homologues the	at function in	Bagellar blögen	hraA	hrc\$	hedT -	hicu	hroV
United	hroC hroJ	hrcN hrpE	hrpQ.		hrpU becs	hrpC hrdT		hrpO (hrpf) hroV (hrp.12) hrpf
R. solensoverum E. emylovoru P. syringse	hroC hroJ hrpH hrpC hrpA1 hrpB3.	hrcN hrp.14 hrp86	hroQ hrpU2/U hrpO1	hrold.	hrpOs hrpOs	hrpX hrpB8 MR	hrpY hrpC1 mB	hrpC2 RhA
X. compostris	niper.	#	BY,N					

a. Gougn et al., 1984, 1985, bid; Wel and Beer, 1983, Bid; 6: V. Beer, unpublished.
b. Bogdanove et al., 1995, bid; Wel and Beer, 1983, Bid; 6: V. Beer, unpublished.
c. Huang et al., 1992, bid; Huang et al., Mol Plant-Microbe Interact 6: 515-520, 1993; Huang et al., 1995, bid; Lidell and Hutcheson, Mol Plant-Microbe Interact 6: 515-520, 1993; Huang et al., 1995, bid; Lidell and Hutcheson, Mol Plant-Microbe Interact 6: 515-520, 1993; Huang et al., 1995, bid; Lidell and Hutcheson, Mol Plant-Microbe Interact 6: 515-520, 1993; Huang et al., 1995, bid; Lidell and Hutcheson, Mol Plant-Microbe Interact 6: 515-520, 1993; Huang et al., 1995, bid; Lidell and Hutcheson, Mol Plant-Microbe Interact 6: 515-520, 1993; Huang et al., 1995, bid; Huang et e. riusing or as, 1864, and 1864, preston of al., 1995, ibid. The predicted product of http:// aligns with the N-terminal two-thirds of a multiple ANGRODE STRETECT (1 200-201) 100-1 1

genes will be designated (#WA, and #WA and Bonas, 1995, bld.; U. Bonas, unpublished. Hwang et al. (1982, J Bacteriof 174; 1923-1831) published d. Fenselau et al., 1992, bld.; Fenselau and Bonas, 1995, bld.; U. Bonas, unpublished. Hwang et al. (1982, J Bacteriof 174; 1923-1831) published c. rensembler su, 1982, but, response and bullet, 1990, but, c. porms, improved the request of su, 1982 and happy such as the sequence of two genes from Xanthomoras campostris pv, glycines, designated 'ORF1' and 'ORF2, that are homologous to happy and happy of

X. campastris pv. vesicatoria, respectively. e. For leviews, see Blair (1995, Bid.) and Electroil and Ordal (1992, Eld.).

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may grow. For any new hip genes that may be discovered, we recommend the strict, sequence-alignment-based criterion for use of the 'hiro' designation until sufficient structural and functional studies can be completed.

Some hip genes are conserved only within subgroups of plant pathogens. One example is the regulatory gene hrpB of R. solana cearum (Genin et al., 1992, Ibid.). This gene, a member of the araC family, is present also in pathovars of X. campestris (Kamdar et al., 1993, J Bacteriol 175; 2017-2025; Karnoum and Kado, 1990, J Bacteriol 172: 5165-5172; U. Bonas, unpublished), but absent from the hrp gene clusters of P. syringee and E. amylovora, which contain regulatory genes that are members of the twocomponent regulatory-system family (Grimm et al., 1995, Mol Microbiol 15: 155-165; Grimm and Panopoulos, 1989, J Bacteriol 171: 5031-5038; XIao et al. 1994, ibid.; S. V. Beer, unpublished). As another example, the hrp gene clusters of P. syringse and E. smylovora each contain a homologue of the Yersinia gene yoph (Bogdanove et al., 1996, ibid.), yet no homologue of this gene has been found in R. solanaceanim or X. campastria. It is noteworthy that the genetic organizations of the hrp gene clusters of X. campestris and R. solanacearum are quite similar to, yet distinct from, those of P. syringee and E amylovora, which resemble one another. We will not attempt a nomenciatural revision here for any of the non-hiro genes, but we encourage authors, wherever possible, to standardize names for such genes, at least within these subgroups, by using conventional rules for bacterial genetic nomenclature, including priority of publication, as a basis for naming homologues (Demerec et al. 1968, Genetics 54: 61-76). Although the same name might be used for different genes across subgroups, startdardized names and the similar genetic organizations within the subgroups should greatly facilitate comparative studies and application of information learned in one species to the study of another.

As for the definition of the 'hrp' designation, it now may include not only genes with a Hrp phenotype, but any gene associated with the Hrp pathway by function, homology, or location within a gene cluster or operon that is essential for the Hrp phenotype. We view use of the 'hrp' designation in this larger sense as elective rather than mandatory. For example, the designation 'hpe' has been used for Hipassociated genes shown not to have a strict Hrp phenotype in R. scianacearum (Gough et al., 1993, ibid.). in order to minimize confusion in the literature, we propose that this designation be maintained for such genes in this organism and in X. campestris, However, for P. syringae and the erwinize, in which gene phenotypes may differ from species to species, we propose a unified nomenclature based on the more inclusive definition of hrp genes presented here. We hope that this broadened definition will help us to gain a focussed understanding of the key

elements underlying the varied and intricate interactions of bacteria with plants.

For convenience, and because 'hrc' represents a subset of hrp genes, hrc and hrp genes collectively will be referred to in general discussion as 'hrp', as in the phrase 'the hrp genes of phytopathogenic bacteria." The combined designation 'hrp/c' may be used to specify a small group of genes, e.g. 'The genes are arranged co-linearly with their hrp/c homologues in Xanthomonas campestris pv. vesicatoria." Operons containing hrc genes still may be referred to as 'hrp' operons. When discussing homologues with the same name (hrp or 'hrc) from more than one plant pathogen, distinctions can be made where necessary using abbreviations for the names of the different bacteria subscripted to the gene name.

The unified nomenclature for conserved hip genes will benefit research in several ways, it makes the known homologies among plant pathogens explicit. It provides for easy cross-reference to other systems, particulary that of Yersinis spp. It facilitates writing and speaking cogently about hip genes. Finally, it transforms a previously confusing jumble of gene names into a well-ordered catalogue, which is an accessible reference not only for hip researchers, but also for those studying other type III secretion systems.

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Regulation of *hrp* Genes and Type III Protein Secretion in *Erwinia amylovora* by HrpX/HrpY, a Novel Two-Component System, and HrpS

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Two novel regulatory components, hrpX and hrpY, of the hrp system of Erwinia amylovora were identified. The hrpXY operon is expressed in rich media, but its transcription is increased threefold by low pH, nutrient, and temperature levels-conditions that mimic the plant apoplast. hrpXY is autoregulated and directs the expression of hrpL; hrpL, in turn, activates transcription of other loci in the hrp gene cluster (Z.-M. Wei and S. V. Beer, J. Bacteriol. 177:6201-6210, 1995). The deduced amino -acid sequences of hrpX and hrpY are similar to bacterial two-component VsrA/VsrD of Pseudomonas regulators including (Ralstonia) solanacearum, DegS/DegU of Bacillus subtilis, and UhpB/UhpA and NarX/NarP, NarL of Escherichia coli. The N-terminal signal-input domain of HrpX contains PAS domain repeats. hrpS, located downstream of hrpXY, encodes a protein with homology to WtsA (HrpS) of Erwinia (Pantoea) stewartii, HrpR and HrpS of Pseudomonas syringae, and other os4-dependent, enhancerbinding proteins. Transcription of hrpS also is induced under conditions that mimic the plant apoplast. However, hrpS is not autoregulated, and its expression is not affected by hrpXY. When hrpS or hrpL were provided on multicopy plasmids, both hrpX and hrpY mutants recovered the ability to elicit the hypersensitive reaction in tobacco. This confirms that hrpS and hrpL are not epistatic to hrpXY. A model of the regulatory cascades leading to the induction of the E. amylovora type III system is proposed.

Additional keywords: fire blight, pathogenicity, virulence.

Erwinia amylovora is the causal agent of the fire blight disease of many rosaceous plants including pear and apple (van der Zwet and Beer 1999). The bacterium infects blossoms, leaves, succulent shoots, and immature fruits. Symptoms of the infected plants include water soaking and discoloration,

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followed by necrosis. Sometimes the disease kills whole trees or substantial portions, resulting in devastating economic loss. In nonhost plants such as tobacco and Arabidopsis, the bacterium elicits the defensive hypersensitive reaction (HR), which is characterized by rapid, localized, cell death (Goodman and Novacky 1994). For infection and HR induction, genes generally called hrp (hypersensitive response and pathogenicity; see Alfano and Collmer 1996 for a review) are essential.

The hrp gene cluster of E. amylovora Ea321 has been cloned in several cosmids and enables nonpathogenic bacteria such as Escherichia coli to elicit the HR in plants (Beer et al. 1991). According to phenotypic analyses of mutants, hrp genes of E. amylovora are localized within a 25-kb region of DNA, consisting of at least eight transcriptional units (Wei and Beer 1993). Sequence analysis (Bogdanove et al. 1996; Kim et al. 1997) indicated that the majority of hrp genes encode proteins that are thought to be components of a specialized protein secretion apparatus called the type III pathway (Hrp pathway for plant pathogens) (Galán and Bliska 1996). Several proteins including harpins (HrpN and HrpW) and a pathogenicity/avirulence protein (DspE) have been shown to be secreted via the pathway (Bogdanove et al. 1998a; Kim and Beer 1998; Wei and Beer 1993).

Transcriptional expression of hrp genes is induced under conditions similar to the environment of the plant apoplast: low carbon and nitrogen, low pH (5.5), and low temperature (18°C) (Wei et al. 1992). Two independent loci, complementation groups IV and V, in the hrp cluster were found to have regulatory function (Sneath et al. 1990; Wei and Beer 1993, 1995). Mutations in these loci abolish harpin production and the HR-eliciting and disease-causing abilities of E. amylovora (Wei and Beer 1993). Preliminary sequence analysis indicated that one of them (group IV) contains a gene called hrpS (Sneath et al. 1990) that encodes a protein similar to 634. dependent transcriptional activators (Morett and Segovia 1993). Complementation group V encodes hrpL (Wei and Beer 1995), which is homologous to genes encoding members of the ECF subfamily of eubacterial sigma factors (Lonetto et al. 1994). HrpL recognizes conserved promoter sequences called "hrp boxes" (Xiao and Hutcheson 1994), and directs the transcription of other pathogenicity genes including hrp secretion operons (hrpA, hrpC, and hrpJ) (Wei and Beer 1995), harpin genes (hrpN and hrpW) (Kim and Beer 1998; Wei and Beer 1995), and a disease-specific locus (dspEF [Bogdanove et al. 1998b]; dspAB [Gaudriault et al. 1997]).

Here we report the characterization of two new regulatory genes, designated hrpX and hrpY, and the further analysis of hrpS. hrpX and hrpY are present in an operon situated between hrpS and hrpL. Analysis of deduced protein sequences suggested that they constitute a two-component regulatory complex; HrpX functioning as a sensor and HrpY as the response-regulator partner of HrpX. hrpX, hrpY, and hrpS are components of a complex regulatory network that leads to activation of hrpL and eventually other genes in the hrp cluster of E. amylovora.

RESULTS

Identification and sequence analysis of the hrpXY locus.

Previous studies have identified several loci, including hrpC, hrpA, hrpS, hrpL, and hrpI, that are essential for the Hrp phenotype (Bogdanove et al. 1996; Kim et al. 1997; Wei and Beer 1993, 1995) (Fig. 1A). Preliminary genetic analysis of pCPP430 in Escherichia coli suggested the presence of a new locus, between hrpS and hrpL, that also is required for the Hrp phenotype and contains novel regulatory components. We have designated this locus hrpXY.

A 3.4-kb Bg/II- and ClaI-digested fragment of pCPP430 was cloned into pBluescript KS+, resulting in pCPP1178. The sequence of the insert of pCPP1178 revealed two tightly linked open reading frames (ORFs) between hrpL and hrpS that are capable of encoding proteins of 495 and 213 amino acid residues, respectively (Fig. 1B). These ORFs were named hrpX and hrpY, respectively. Potential ribosome-binding sites, AGGAG and TGGAA, were found 5 and 7 bp upstream of the hrpX and hrpY start codons, respectively. Although the ribosome-binding site ahead of hrpY weakly matches the consensus sequence, we assume it is sufficient for translation of hrpY; only a 4-bp space exists between the hrpX stop codon and hrpY start codon and translational coupling is plausible. To confirm that

the hrpX and hrpY ORFs produce proteins, pCPP1178 was placed in a gene expression system mediated by the T7 RNA polymerase. Two distinct protein bands were visible following sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). The apparent molecular masses of HrpX and HrpY were about 50 and 25 kDa, respectively (data not shown), close to the sizes expected from the deduced amino acid sequences.

The start codon of hrpX is located 146 bp downstream of the hrpL stop codon, and a promoter prediction program (see Materials and Methods) identified two putative on promoter sequences, TAGACG-N_{IT}TAAAGT (score from promoter prediction by neural network = 0.97) and TTGCAA-N_{IS}-CCTAAT (score = 0.95), 111 and 33 bp upstream of the hrpX start codon, respectively. There is a 361-bp noncoding region between hrpY and hrpS. Palindromic sequences that may serve either as targets of regulatory components or as transcription terminators, GTAAACANTGTTTAC and GGATAAAATGG-TTGTGG-N_T-CCGCTTCCATTTTATCC, were identified in the hrpL-hrpX and hrpY-hrpS intergenic regions, respectively. The tight linkage of hrpX and hrpY, and the existence of long noncoding areas and inverted repeats upstream of hrpX and downstream of hrpY, suggest that the two genes form an operon.

HrpX and HrpY constitute

a two-component regulatory system.

Comparison of the predicted amino acid sequences of hrpX and hrpY with sequences in the data bases revealed significant similarities with many two-component regulatory proteins. The homologs include VsrA/VsrD of Pseudomonas (now Ralstonia) solanacearum, which regulate virulence gene expression (Huang et al. 1995b); UhpB/UhpA of Escherichia coll, which participate in the regulation of sugar transport (Friedrich and Kadner 1987); NarX/NarP,NarL of Escherichia

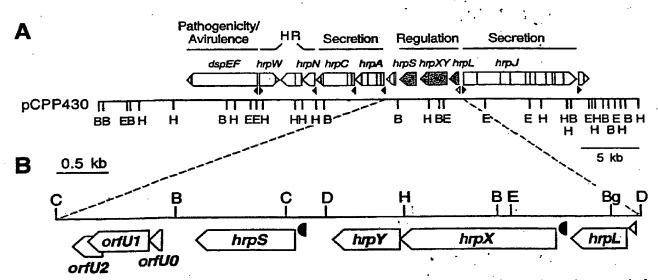


Fig. 1. A, Operon organization of the hrpldsp gene cluster of Erwinia amylovora cloned in pCPP430. B, Central region covering regulatory genes hrpl., hrpl., and hrpl. Boxes and arrow boxes: transcriptional units or open reading frames; names of the characterized operons or genes are given above, inside, or below. Filled triangles: putative Hrpl.-dependent promoters. Open triangles: putative of promoters. Closed half circles: putative of promoters. Restriction enzymes: B, BamH; E, EcoRI; H, HindIII; Bg, BgIII; C, ClaI; and D, DraI.

coll, which are involved in the regulation of anaerobic respiratory gene expression (Rabin and Stewart 1993); and DegS/DegU of Bacillus subtilis, which are involved in extracellular enzyme production (Kunst et al. 1988) (Fig. 2; Table 1). In addition, HrpY showed high sequence similarity with many other transcriptional activators including ExpA of E. carotovora (33% identity), which is involved in global control

of virulence (Eriksson et al. 1998); UvrY of Escherichia coli (33% identity) (Sharma et al. 1986); SirA of Salmonella typhimurium (32% identity) (Iohnston et al. 1996); and GacA of several animal- and plant-associated Pseudomonas spp. (29 to 30% identities) (Laville et al. 1992).

The high sequence similarity of HrpX with histidine kinases suggests that HrpX is a sensor. HrpX has the conserved His

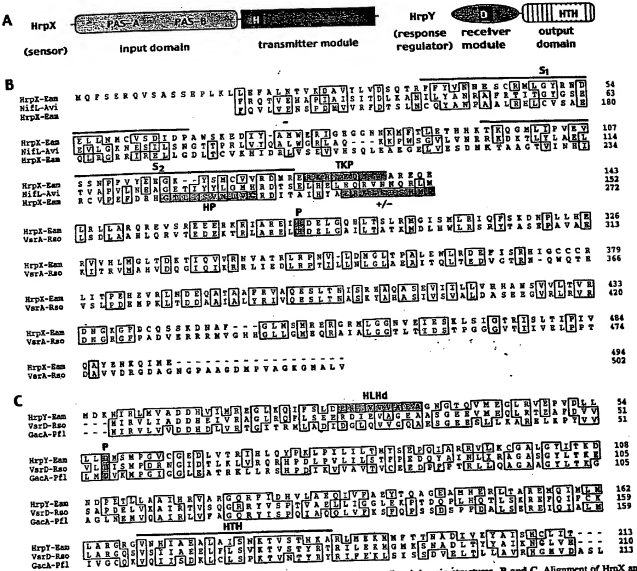


Fig. 2. A, Diagramatic illustrations of HrpX and HrpY of Erwinia amylovora showing predicted domain structures. B and C, Alignment of HrpX and HrpY with similar proteins. Designations of diagrams are after Parkinson and Kofoid (1992). PAS-A and PAS-B denote two repeats of the PAS domain, HrpY with similar proteins. Designations of diagrams are after Parkinson and Kofoid (1992). PAS-A and PAS-B denote two repeats of the PAS domain, Government and aspartate residues, respectively, and HTH the helix-num-helix DNA-binding motif. Overlines represent the S-boxes in the PAS domain (S₁ and S₂) and the HTH motif. A putative tyrosine kinase phosphorylation site (TKP), a hydrophobic region (HP), a putative charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and a helix-loop-helix dimerization domain signature (HLHd), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and phosphorylation sites (P) are shown by shading. PILEUP charge-rich linker (4/-), and phosphorylation sites (P) are shown by shading rich shading rich shading rich shading rich shading rich shading ric

residue for autophosphorylation and a hydrophobic domain that may enable the protein to be transiently associated with the cytoplasmic membrane (Fig. 2B). The C-terminal putative transmitter domain (residues 273 to 494) of HrpX shows most similarity to the kinase domains of the sensor proteins listed in Table 1; the N-terminal putative input domain of HrpX shows similarity to PAS domains (Zhulin et al. 1997) of Methanobacterium thermoautotrophicum, Azotobacter vinelandii, and other organisms. Several PAS-containing proteins are sensors of bacterial two-component systems. The PAS domain typically consists of two direct sequence repeats (PAS-A and PAS-B), and each repeat contains two highly conserved regions called S1 and S2 boxes (Zhulin et al. 1997). In the case of HrpX, the second repeat (PAS-B) seems imperfect (Fig. 2B). Based on ScanProsite analysis (Appel et al. 1994), another feature of HrpX with unknown functional relevance is a putative tyrosine kinase phosphorylation site (PROSITE:PS00007).

HrpY appears to be a response regulator with a putative receiver domain at the N terminus (up to 102 amino acid residues) and a DNA-binding domain at the C terminus (Fig. 2A). As shown in Figure 2C, HrpY contains the conserved Asp residue, which may be phosphorylated by the sensor, and the

Table 1. HrpX and HrpY of Erwinia amylovora compared with twocomponent regulatory proteins (sensors/response regulators) of other

Becterium	Protein	Amino ecids	% Identity
Erwinia amylovora Ralstonia solanacearum Escherichia coli Bacillus subtilis Escherichia coli	HrpX/HrpY VsrA/VsrD UhpB/UhpA DegS/DegU NarX/NarP, NarL	494/213 502/210 500/196 385/229 598/215, 216	34/41 32/32 32/28 31/33, 32

6 % Identities from a BLASTP search of HrpX and HrpY with default parameters, except for no filtering for low complexicity regions. Only the transmitter domain of HrpX (residues 273 to 494) was used for comparisons with other sensor proteins.

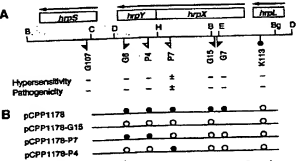


Fig. 3. Genetic characterization of the hrpXY locus. A, Locations of transposon insertions and phenotypes of hrpX and hrpY mutants of Erwinia amylovora Ea321. Rectangles above map of restriction enzymes and transposons represent transcriptional units. Arrows: directions of transcription. Closed flags: insertions by Tn5-gusA1. Open flags: insertions by TnphoA. Llollipop: a Tn10-miniKm insertion. Mutants shown by minus signs below insertion points did not elicit the hypersensitive reaction (HR) or cause disease (Hrp*); a mutant shown by ± infrequently elicited spotty HR and showed low virulence. B, Complementation assay of hrpX and hrpY mutants of E. amylovora Ea321 with various plasmids. Closed circle: plasmid complemented Hrp phenotype of the mutant containing the transposon insertion in the same column. Open circle: plasmid did not change the phenotype of corresponding mutant.

helix-turn-helix DNA binding motif. HrpY also has a sequence that matches the Myc-type helix-loop-helix dimerization domain signature (PROSITE-PS00038), the functional significance of which remains to be tested.

Genetic characterization of hrpX and hrpY.

The hrpXY locus in pCPP430 was mutagenized with transposons Tn5-gusA1 and TnphoA. Derivatives of pCPP430 containing the transposon insertions were marker-exchanged into the genome of E. amylovora Ea321. All hrpY mutants of Ea321 failed to elicit the HR in tobacco and to infect immature pear fruits (Fig. 3A). Two classes of hrpX insertion mutants were obtained. Ea321-G15 and Ea321-G7, which were made with Tn5-gusA1, were similar to hrpY mutants in phenotypes. Ea321-P7, an hrpX::TnphoA mutant, caused slight tissue collapse in tobacco at higher inoculum dose and had low virulence in immature pears, rather than the strict Hrpphenotype (Fig. 3A). Specifically, tobacco leaves infiltrated with Ea321-P7 at ≥ 5 × 108 CFU per mi developed a spotty HR 36 h after infiltration. Also, in immature pears inoculated with the mutant, bacterial ooze appeared 3 days later than in those inoculated with the wild type, and the population of the mutant recovered was only one-tenth of that of the wild type (data not shown).

Virulence of the mutants was restored to near wild-type levels by providing the mutants with pCPP1178 in trans (Fig. 3B). The hrpX::Tn5-gusAl mutants of Ea321 were not complemented by pCPP1178-P4 that contains a transposon insertion in hrpY (Fig. 3B). This suggests that hrpX and hrpY are in the same transcriptional unit and the Tn5-gusA1 mutations in hrpX are polar. We found, however, that the hrpX::TnphoA mutant Ea321-P7 can be complemented by pCPP1178-P4, indicating that the TnphoA insertion of hrpX did not affect the function of hrpY (Fig. 3B). TnphoA-induced mutations that permit the expression of downstream genes have been observed frequently in E. amylovora (Z. Wei and S. V. Beer, unpublished data) and Pseudomonas syringae (Huang et al. 1995a). Thus, we believe that the P7 insertion is nonpolar and that the peculiar phenotype of the Ea321-P7 may reflect the function of hrpX.

All the transposon mutations in the hrpXY locus were complemented by derivatives of pCPP430 with transposon insertions in hrpS or hrpL (data not shown), confirming the suggestion from sequence analysis that hrpX and hrpY constitute an independent complementation group. Based on results of sequence analysis and genetic characterization, we conclude (i) hrpXY is required for the Hrp phenotype, and (ii) hrpX and hrpY constitute a two-gene operon, hrpXY.

Expression of hrpXY is environmentally regulated.

A new construct, pCPP1203, was used to monitor expression of the hrpXY promoter in a nutrient-rich medium and a minimal medium that induces the expression of hrp genes (Wei et al. 1992). pCPP1203 was derived from pCPP1178-G15 (hrpX::Tn5-gusA1) in which the directions of hrpX and gusA are the same. pCPP1178-G15 was digested with BamHI and SacI (an SacI site is present in the vector), which cuts out the hrpXY promoter region, a 5' portion of the hrpX coding region fused to Tn5-gusA1, and the whole transposon. The resulting fragment was then ligated to pCPP43, which had been digested with the same enzymes. pCPP43 (gift of David

W. Bauer) is a derivative of pOU61, which is a low-copynumber plasmid (approximately one copy per bacterium at 30°C) (Larsen et al. 1984).

In E. amylovora and Escherichia coli, the hrpXY promoter directed high levels of basal expression in Luria broth (LB), but expression of hrpX::Tn5-gusAl was enhanced threefold in the hrp-inducing minimal medium (IM) (Table 2). Enhanced levels of hrpX::Tn3-gusA1 expression were also observed from assays of the strains in tobacco leaves and immature pears (data not shown). No \(\beta\)-glucuronidase (GUS) activity was detected for Escherichia coli SØ200\(DidA(pCPP1203)\) unless functional hrpXY was provided (Table 2). Similarly, high basal-level expression of hrpX::Tn5-gusA1 of Ea321(pCPP1203) in Table 2 is probably due to functional hrpXY present in the chromosome. The latter two observations indicate that hrpXY is also au-

hrpX and hrpY control the expression of hrpL.

To study the effect of hrpX and hrpY on the control of hrpL expression, a hrpL::Tn5-gusAl fusion (pCPP139-G44) (Wei and Beer 1995) was marker exchanged into an hrpX mutant (Ea321-P7) and an hrpY mutant (Ea321-P4), to generate hrpXhrpL and hrpY-hrpL double mutants Ea321-P7G44 and Ea321-P4G44, respectively. Mutation in hrpY completely abolished hrpL expression (Fig. 4). However, the hrpX mutant reduced hrpL expression only to about 20% of its wild-type level, opening the possibility that the mutated HrpX may be still partially functional or another sensor protein may cross talk with HrpY.

Analysis of the hrpS locus and the ORFs between hrpS and hrpA.

hrpS also partially controls hrpL expression in E. amylovora and is located downstream of hrpXY (Wei and Beer 1995). We report here the entire nucleotide sequence of the region between hrpY and hrpA, which includes hrpS, to complete the preliminary results on hrpS presented previously (Sneath et al. 1990).

The hrpS locus of E. amylovora Ea321 contains a single-gene operon, based on the large intergenic regions beyond the coding region of hrpS, and a potential terminator, CGGCGACAGC-Ns-GCTGTCGCCG, that lies 49 bp downstream of the hrpS stop codon. The hrpS ORF is preceded by a potential σ^{70} promoter, GTGGCA-N₁₈-TATTAC (score from promoter prediction by neural network = 0.96), and it encodes a 324 amino acid protein. HrpS has homology to members of the o54-dependent, enhancerbinding protein family (Morett and Segovia 1993). HrpS shows highest sequence similarity with WtsA (HrpS) of Erwinia (Pantoea) stewartii (Frederick et al. 1993) (79% identity over 322 amino acid residues without gaps from BLASTP), HrpR and HrpS of P. syringae pathovars (51 to 55% identities) (Grimm et al. 1995; Xiao et al. 1994), and DctD of Rhizobium spp. (39% identities) (Jiang et al. 1989; Ronson et al. 1987). HrpS of E. amylovora has two putative ATP-binding sites at the N terminus and a belix-turn-helix DNA-binding motif at the C terminus (Fig. 5A). HrpS shows high sequence similarity to other regulators in the NuC family throughout the entire out interaction domain. However, similar to other HrpR/HrpS proteins, HrpS of E. amylovora contains a very short N-terminal A domain (Shingler 1996), and seems to lack the phosphorylation receiver domain (Fig. 5A).

In the region between hrpS and hrpA, three potential genes, designated orfU0, orfU1, and orfU2 (Fig. 1B), were identified by application of the GeneMark.hmmn algorithm (Lukashin and Borodovsky 1998). orfU0 is a small ORF encoding a 46 amino acid basic protein, without significant similarity to any protein in the data base. Preceded by GGAGT 8 bp upstream, orfUI encodes a 203 amino acid basic protein that is similar to a conserved hypothetical protein HP14O1 of Helicobacter pylori (32% identity over 164 amino acid residues with 12 gaps) (Fig. 5B). Interestingly, protein sequence of the next ORF, orfU2, shows even higher similarity to HP1401 (residues 189 to 229; 41% identity without gaps). This suggests the possibility that a frame shift in orfU1-orfU2 resulted in the two current ORFs, and that both may be defective. The lack of an obvious promoter in front of orfUO, the lack of good ribosome-binding sites in front of orfU0 and orfU2, the potential frame-shift mutation at the 3' region of orfU1, and the lack of a phenotype of TnphoAinduced orfUl mutants (data not shown) indicate that the region comprising orfU0-orfU2 is unlikely to be functional in Ea321.

Expression of hrpS is not autoregulated, and induction of hrpS is independent of hrpX or hrpY.

An hrpS::gusAl fusion designated G107 (Wei et al. 1992) was used to assay the expression of hrps. A fragment of

Table 2. Expression of the hrpXY promoter in Luria broth (LB) and in a hrp-inducing minimal medium (IM)

	GUS activity			
Bacterial strains	LB	IM		
Erwinia amylovora Ea321(pCPP1203) E. coli SØ200\(\Delta\text{uidA(pCPP1203)}\) E. coli SØ200\(\Delta\text{uidA(pCPP1203)}\) P. coli SØ200\(\Delta\text{uidA(pCPP1203)}\)	242 ± 12 2 ± 3 145 ± 19	788 ± 32 3 ± 3 878 ± 33		

^{*} E. coli SØ200ΔuidA is an Escherichia coli strain with no β-glucuronidase (GUS) activity due to deletion of gusA. pCPP1203 is a low number plasmid containing hrpX::Tn5-gusAI; pCPP1178 is a high-copy-number plasmid containing-functional hrpX and hrpY genes.

Picounits per CFU; mean of three replicates ± standard deviation.

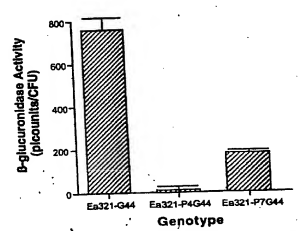


Fig. 4. Effect of mutations in hrpX and hrpY on expression of hrpL. Genotypes of the strains are Ea321-G44, hrpL::Ta5-gusA1 (Wei and Beer 1995); Ea321-P4G44, hrpY::TnphoA and hrpL::Tn5-gusAl; and Ea321-P7G44, hrpX::TnphoA and hrpL::Tn5-gusAI. Error bars: standard deviation from three replicates. Cells grown in inducing medium (IM) were assayed (see Materials and Methods for details).

pCPP430-G107 digested with BamHI contains the whole transposon, the hrpS gene fused to Tn5-gusAI, and the hrpS promoter region. This BamHI fragment was ligated with a low-copy-number plasmid, pCPP8 (Bauer 1990), that was cut with the same enzyme. The resulting plasmid was designated pCPP1058. As with hrpXY, expression of hrpS in Escherichia coli or in E. amylovora was induced under hrp-inducing conditions (Table 3). However, autoregulation was not required for hrpS expression; the presence of functional hrpS did not affect the expression of a hrpS::gusAI fusion in pCPP1058 (Table 3).

To determine whether the newly discovered two-component system has any effect on the expression of hrpS, an hrpS::TnS-gusA1 fusion (pCPP430-G107) was marker-exchanged into hrpX and hrpY mutants. Neither hrpX nor hrpY affected hrpS expression significantly (Fig. 6).

hrpS and hrpL, provided by multicopy plasmids, suppress defects in hrpX or hrpY.

To further characterize the regulatory relationships between hrpXY, hrpS, and hrpL, the HR-impaired strains Ea321-P7, Ea321-P4, and Ea321-G107 were transformed with pCPP1178

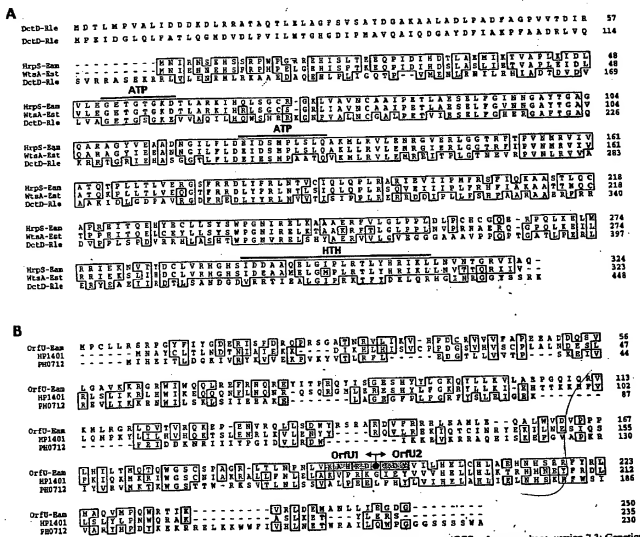


Fig. 5. Alignments of A, HrpS, and B, OrfU of Erwinia amylovora with similar proteins. PILEUP program (GCG software package, version 7.3; Genetics Computer Group, Madison, WI, U.S.A.) with default parameters was used to align the sequences. Overlines represent ATP-binding sites (ATP) and the belix-turn-helix DNA-binding motif (HTH). Sequence of OrfU is a composite of sequences of orfU1 and orfU2 products. A putative tyrosine kinase phosphorylation site (PROSITE: PS00007) is indicated by shading. Black circle in the OrfU sequence denotes location of a probable reading-frame shift. Accession numbers: WtsA of E. stewartii, SWISS-PROT:P36219; DctD of Rhizobium leguminosarium, SWISS-PROT: P10046; HP1401 of Helicobacter pylori, GENBANK:AE000640; and PH0712 of Pyrococcus horikoshil, DDBJ:AP000003.

(contains hrpXY), pCPP1001 (contains hrpS) (Wei and Beer 1995), or pCPP1078 (contains hrpL) (Wei and Beer 1995). The resulting transformants were infiltrated into panels of tobacco leaves to determine which, if any, of the regulatory genes, when present in multiple copies, are sufficient to restore the HR-eliciting ability to the mutants. Panels infiltrated with hrpX and hrpY mutants containing hrpL developed the HR (Table 4), often faster than panels infiltrated with the wildtype strain. The panels began to show collapse 8 to 12 h after infiltration; by 24 h, the whole infiltrated area had collapsed in a typical HR. This result is consistent with dependence of hrpL expression on hrpX and hrpY. Interestingly, similar suppression was observed from hrpX and hrpY mutants containing hrpS, whereas hrpX and hrpY did not restore the HR phenotype of the hrpS mutant (Table 4).

DISCUSSION

The HrpX/HrpY two-component protein system.

Our results demonstrate that E. amylovora employs the HrpX/HrpY two-component regulatory proteins to direct expression of an alternate sigma factor gene, hrpL, that in turn activates a type III protein secretion system. This provides for a quick change in the pattern of gene expression needed to initiate infection. HrpX is a putative IcT-type sensor (Parkinson and Kofoid 1992) composed of the N-terminal PAS domain and the C-terminal histidine kinase domain (Fig. 2A). HrpX appears to be cytoplasmic, and may be anchored to the inner membrane by its internal hydrophobic region. Other members of the PAS-containing LeT-type sensor kinases include Nifl., NtrB, and KinA (Zhulin et al. 1997). HrpY appears to be a ROm subfamily response regulator (Parkinson and Kofoid 1992). Consistent with the HrpX transmitter domain, HrpY shows significant sequence similarity to VsrD, DegU, UhpA, and NarL.

Two-component systems with PAS domains in the sensor component include NifL/NifA, DctS/DctR, and BvgS/BvgA (Zhulin et al. 1997). Among these only NifL does not contain the periplasmic domain, and HrpX is more similar to NifL than the other two. NifL and most other PAS-containing proteins are sensors (Zhulin et al. 1997), and their signal input domains are located at the N terminus (Parkinson and Kofoid 1992). Thus, HrpX may directly perceive environmental signals with its N-terminal PAS domain. One function of the PAS domain is to act as a protein dimerization motif (Kay 1997). This raises the possibility of HrpX dimerization, which is required for the functional state of two-component sensors (Parkinson and Kofoid 1992).

Two-component regulatory system and type III protein secretion.

Although the two-component system is widely used to control bacterial gene expression (Hoch and Silhavy 1995), reports of its function in regulation of the type III system are just emerging. In S. typhimurium, SirA is a response regulator essential for induction of hild, prgHIJK, and sigDE (Hong and Miller 1998; Johnston et al. 1996), and the PhoQ/PhoP twocomponent system represses the expression of the prg locus (Pegues et al. 1995). The CpxA/CpxR system controls the pHdependent expression of the Shigella sonnei virF gene, which in turn activates ipaBCD and virG (Nakayama and Watanabe

Table 3. Expression of the hrpS promoter in Luria broth (LB) and in hrp-inducing minimal medium (IM)

	GUS activity		
Bacterial strain	LB	IM	
E coli SØ200\(\textit{LuidA}\)(pCPP1058) E coli SØ200\(\textit{LuidA}\)(pCPP1058, pCPP1001) Erwinia amylovora Ea321-G107 Erwinia amylovora Ea321-G107(pCPP1001)	94 ± 12 105 ± 17 36 ± 11 42 ± 21	367 ± 9 378 ± 23 188 ± 35 229 ± 29	

*E. coli SØ200ΔuidA is an Escherichia coli strain with no βglucuronidase (GUS) activity due to deletion of gusA. Erwinia amylovora Ea321-G107 is a mutant of Ea321 containing a Ta3-gusAI insertion in htps (Wei et al. 1992). pCPP1058 is a low-copy-number plasmid containing htpX::Tn5-gusAI; pCPP1001 is a high-copy-number plasmid containing the functional hrps gene and its promoter (Wei and

b Picounits per CFU; meanof three replicates ± standard deviation.

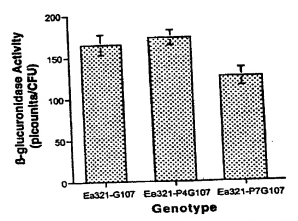


Fig. 6. Effect of mutations in hrpX and hrpY on expression of hrpS. Genotypes of the strains are Ea321-G107, hrpS::Tn5-gusA1 (Wei et al. 1992); Ea321-P4G107, hrpY::TaphoA and hrpS::Ta5-gusAI; and Ea321-P7G107, hrpX::TuphoA and hrpS::Tn5-gusAI. Error bars: standard deviation for three replicates. Cells grown in inducing medium (IM) were assayed (Materials and Methods contains details).

Table 4. Hypersensitive reaction (HR) elicitation by hrp regulation mutants

Strain	Genotype	HR phenotype	
Ea321	wild type; hrp*	+++	
Ea321-P7 Ea321-P7(pCPP1178) Ea321-P7(pCPP1001) Ea321-P7(pCPP1078)	hrpX hrpX(hrpXY*) hrpX(hrpS*) hrpX(hrpL*)	± ++> +++	
Ea321-P4 Ea321-P4(pCPP1178) Ea321-P4(pCPP1001) Ea321-P4(pCPP1078)	hrpY hrpY(hrpXY*) hrpY(hrpS*) hrpY(hrpL*)	++ > +++ +++	
Ea321-G107 Ea321-G107(pCPP1178) Ea321-G107(pCPP1001)	hrp\$ hrp\$(hrpXY*) hrp\$(hrp\$*)	<u>-</u>	

+++, full HR manifested by complete tissue collapse throughout infiltrated area; ++, reduced HR, which is sporty and often coalescing; ±, infrequent collapse and small spotty necreosis for HR-positive leaves; and –, no HR. Inoculum concentration was approximately 2×10^6 CFU per ml. Ratings (consensus of four plants) were made 36 h after in-

^b Full HR was observed at inoculum levels of $\geq 5 \times 10^8$ CFU per ml.

1995). Also, the BvgS/BvgA system was recently found to be involved in the regulation of the type III secretion in Bordetella bronchiseptica (Yuk et al. 1998). Among plant pathogens, HrpG of Xanthomonas campestris pv. vesicatoria, a homolog of response regulators, has been shown to regulate hrpXv and hrpA expression (Wengelnik et al. 1996).

The structure of the input domain of E. amylovora HrpX appears to be exceptional, compared with sensor proteins involved in other type III systems, which contain two transmembrane regions and a periplasmic domain. The closest homologs of E. amylovora HrpY are SirA and BvgA, both of which are RO_{III}-type regulators (Parkinson and Kofoid 1992), whereas X. campestris HrpG belongs to the RO_{II} type, which includes Escherichia coli CpxR and OmpR, S. ryphimurium PhoP, and Agrobacterium tumefaciens VirG. Thus, at least two types of transmitter-receiver systems appear to have evolved for control of type III systems in response to environmental stimuli in hosts. Also, the two two-component systems identified in the plant pathogens E. amylovora and X. campestris fall into different communication groups.

HrpS and mechanism of gene regulation.

HrpS is a member of the o⁵⁴-dependent, enhancer-binding protein family. Both hrpS and rpoN are required for transcrip-

tion of hrp genes in P. syringae pathovars (Grimm et al. 1995; Xiao et al. 1994). WtsA (HrpS) of E. stewariii controls expression of wtsB, which also requires the presence of of (Frederick et al. 1993). In E. amylovora, HrpS partially regulates hrpL expression (Wei and Beer 1995), and a sequence, TGGCAC-N₅-TTGC, that perfectly matches the -241-12 promoter consensus sequence is found at the promoter region of E. amylovora hrpL. The hrpS gene of E. amylovora, but not hrpS of P. syringae pv. phaseolicola, can complement the hrpS mutation in E. stewariii (Frederick et al. 1993). The HrpS sequences of the two erwinias are highly similar, and even the upstream noncoding regions appear to be conserved, except for the insertion of a 484-bp sequence, reminiscent of an IS (insertion sequence) element, 23-bp upstream of the E. stewariii hrpS ORF.

As a member of the NtrC family, HrpS is unusual in that it lacks a long N-terminal receiver domain. Control of protein activation by phosphorylation, by protein-protein interaction, and by signal molecule have been suggested for o⁵⁴-dependent proteins (Shingler 1996). In the direct activation model, derepression by effectors seems to be a mechanism of protein activation. For DctD, DmpR, and XyIR, deletion of the receiver domain results in constitutive activation of the proteins, suggesting that the receiver domain has a repressor function

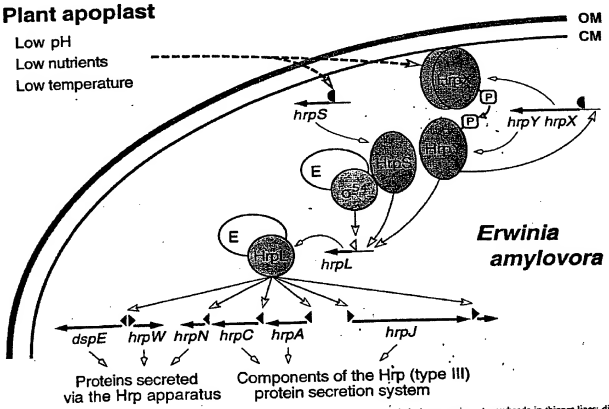


Fig. 7. Model of the hrp gene regulatory cascade. Thick arrow lines: genes or operons. Ovals and circles: proteins. Arrowheads in thinner lines: directions of information flow. CM, cytoplasmic membrane; OM, outer membrane; P, phosphate; E, RNA polymerase; closed half circle, o⁷⁶ promoter; open triangle, o⁷⁶ promoter; and filled triangle, HrpL promoter.

(Shingler 1996). Therefore, the apparent absence of the receiver domain in HrpS implies that HrpS may not require phosphorylation for activation and is always active once the protein is made.

Induction of hrpXY and hrpS and the involvement of HrpXY and HrpS in hrpL regulation.

Expression of hrpS and hrpXY is induced by conditions that mimic the apoplastic environment (Wei et al. 1992; this work). hrpXY shows high basal-level expression, and autoregulation is involved in gene induction. However, hrpS is not autoregulated based on results of the GUS assay, suggesting that there may be upstream regulatory components. Although hrpS provided in multiple-copy plasmids reverses the Hrp phenotype of hrpX and hrpY mutants, the independence of hrpS from hrpX and hrpY suggests that hrpXY is not epistatic to hrpS and environmental signals may go to hrpS through a different pathway.

Earlier work on hrpL and hrpS (Wei and Beer 1995) established that HrpS partially controls hrpL expression. Our current work indicates that the HrpX/HrpY system contributes to hrpL induction. Based on the role of hrpXY and hrpS in regulating hrpL and the lack of effect of hrpX and hrpY in hrpS expression, one might place hrpS upstream of hrpXY. This notion is precluded, however, because hrpXY does not override hrpS mutation. As mentioned above, the opposite is not likely, either. Therefore, it seems that signals independently perceived by hrpXY and hrpS converge at hrpL.

Neither HrpS nor HrpY alone induce high levels of hrpL expression, suggesting that cooperation of HrpY and HrpS, possibly through protein-protein interaction, may be needed for full activation of hrpL. In this model, HrpS may be a positive activator of hrpL, while HrpX/HrpY may act as a modulator of hrpL transcription. Complementation of hrpX and hrpY mutants for the HR phenotype by overexpressed hrpS supports this model. The regulation of eps genes of R solanacearum seems similar, both VsrD and PhcA regulators bind to the xpsR promoter region and control xpsR expression (Huang et al. 1995b). In P. syringae pv. syringae, HrpR and HrpS have been proposed to work together to control hrpL expression (Xiao et al. 1994), although a different opinion exists for homologous proteins in P. syringae pv. phaseolicola (Grimm et al. 1995).

hrp gene regulation and Hrp phenotypes.

hrpY and hrpS seem to be crucial to the pathogenic life-style of E. amylovora, since their inactivation by mutagenesis results in loss of pathogenicity in immature pears (Wei et al. 1992; this work). The hrpX mutant, however, shows an attenuated phenotype: slightly lowered hrpL expression and reduced HR and virulence at higher inoculum doses. Currently, we cannot rule out the possibility of partial HrpX function in that mutant, even though leaky phenotypes of sensor mutants have been documented for other two-component systems (Stock et al. 1989). It is interesting to note that, although hrpX and hrpS mutants show different phenotypes (the former reduced Hrp and the latter Hrp), both are similarly affected in hrpL expression, i.e., only three- to fourfold reduction. This suggests that either there is a threshold level of hrp gene expression required for causing disease, or hrpS is involved in expression of other genes that contribute to pathogenicity. Further study might distinguish between these two possibilities.

The incomplete complementation of hrpX and hrpY mutants by hrpXY provided in a multicopy plasmid at lower inoculum levels ($\leq 2 \times 10^8$ CFU per mI) is intriguing and deserves further investigation. One explanation for the results could be that defective HrpX and HrpY in the mutants interact with functional HrpX and HrpY, and, possibly by forming heterodimers, interfere with the full activity. Alternatively, overproduced HrpX and HrpY may somehow down-regulate hrpS expression.

Model of the E. amylovora hrp gene expression.

Based on previous studies (Bogdanove et al. 1996, 1998b; Kim and Beer 1998; Kim et al. 1997; Wei and Beer 1995; Wei et al. 1992) and results described in this work, we propose a scheme of hrp gene regulation in E. amylovora (Fig. 7). When the bacteria enter the plant apoplast, HrpX perceives environmental signals and is phosphorylated. Activated HrpX then phosphorylates HrpY to activate it, and increases the expression of hrpXY to produce more HrpX and HrpY. Independently, expression of hrpS is induced in response to the changed environment. Activated HrpY and HrpS, bound to the hrpL promoter, then interact with the RNA polymerase-o's complex to drive transcription of hrpL. HrpS also activates other genes containing the -24/-12 promoter consensus sequence. Finally, the HrpL o factor, which recognizes a conserved promoter motif, GGAACC-Nir-CCACTAAT, directs transcription of the remaining hrp and dsp genes that produce the secretion machinery and virulence proteins that interact with plant cells.

MATERIALS AND METHODS

Bacterial strains and growth condition.

E. amylovora Ea321 is a wild-type strain that infects pear and apple (Beer et al. 1991). Escherichia coli DH5α was routinely used for cloning of cosmids and plasmids. pCPP1001 (Wei and Beer 1995), pCPP1036 (Kim et al. 1997), pCPP1078 (Wei and Beer 1995), and pCPP1178 are subclones of pCPP430 (Beer et al. 1991), and contain ORFs in the same direction as the T7Φ10 promoter from the vector pBluescript KS+. Strains of E. amylovora Ea321 and Escherichia coli were grown in LB (Sambrook et al. 1989) with vigorous shaking at 28 and 37°C, respectively. Inducing medium (IM) was used for inducing hrp gene expression as described previously (Wei et al. 1992). The antibiotics used to maintain selection were ampicillin at 100 μg/ml, kanamycin (Km) at 50 μg/ml, spectinomycin (Sp) at 50 μg/ml, tetracycline (Tc) at 20 μg/ml, and carbenicillin (Cb) at 300 μg/ml.

Recombinant DNA techniques and sequence analysis.

Unless otherwise specified, basic molecular biology techniques were as described (Sambrook et al. 1989). Electroporation of plasmid DNA into *E. amylovora* 321 and its derivatives was done as described by Bauer and Beer (1991) with the Gene Pulser apparatus (Bio-Rad, Richmond, CA, U.S.A.).

Deletion clones, generated from the ClaI-BgIII insert in pCPP1178 with the Erase-A-Base kit (Promega, Madison, WI, U.S.A.), were sequenced by the dideoxy chain termination procedure with the Sequenase sequencing kit (U.S. Biochemical, Cleveland, OH, U.S.A.). Also, sequencing of the region between hrpA and hrpJ in pCPP430, pCPP1001, pCPP1036, and pCPP1178 was performed on an ABI 373A automated DNA sequencer (Perkin-Elmer, Norwalk, CT, U.S.A.) at the

Cornell University Biotechnology Program DNA Sequencing Facility with oligonucleotide primers synthesized at the same

DNA and deduced amino acid sequences were analyzed with programs in the GCG software package, version 7.3 (Genetics Computer Group, Madison, WI, U.S.A.) and DNASTAR (DNASTAR, Madison, WI, U.S.A.). Potential genes were identified with GeneMark.hmm (Lukashin and Borodovsky 1998; available on-line from the GeneMark web site). Homology searches were done with BLAST algorithms (Altschul et al. 1997; available on-line from the NCBI web site). Conserved patterns in proteins were found with Scan-Prosite (Appel et al. 1994; available on-line). Finally, prediction of potential o70 promoters were made with the Promoter Prediction by Neural Network method (Reese and Eeckman 1995; available on-line).

Expression of hrpX and hrpY in Escherichia coli.

A gene expression system mediated by a T7 RNA polymerase/promoter (Tabor and Richardson 1985) was used. pCPP1178, which contains hrpX and hrpY ORFs driven by the T7Φ10 promoter from the vector, was introduced into Escherichia coli DH5cu(pGP1-2). Cells were incubated at 42°C to induce the expression of the T7 RNA polymerase gene, and newly synthesized proteins were radiolabeled with ³⁵S-Met as described (Tabor and Richardson 1985). Resulting samples were resuspended in a sample buffer and heated to 95°C for 3 min before being electrophoresed in a 12% polyacrylamide gel.

Construction of marker-exchange mutants.

Chromosomal mutants were constructed by markerexchange mutagenesis as described previously (Wei et al. 1992). A Tulo-minikm insertion or a TuphoA insertion, mapped at the hrpXY or hrpL locus in Escherichia coli DH5(pCCPP430) or Escherichia coli DH5α(pCPP1178), was introduced into E. amylovora Ea321 by triparental mating with the helper strain, Escherichia coli HB101(pRK600) (kindly provided by E. R. Signer; Department of Biology, Massachusetts Institute of Technology, Cambridge). The transconjugants were selected on Luria plates containing Km and Sp, and then transferred to a low-phosphate minimal medium (Bauer 1990) to select for Kmr Spr marker-exchanged mutants. The second mutations were generated by introducing individual hrp::Tn5-gusAl fusions into Tn10-miniKm or TnphoA mutants of Ea321. Since the transposon Tn5-gusAl has two selection marker, Km and Tc, the second mutation was selected based on Km' Te' Sp' phenotype. All the mutants were tested for the HR-eliciting ability and pathogenicity. TnphoA insertions P74 and P86 in pCPP1036, which were mapped to orfUl, were introduced to the Ea321 genome by electroporation and subsequent incubation in a low-phosphate medium with Km. Integration of the TnphaA fusion into the chromosome was confirmed by antibiotic resistance (Kmr Cbr) and Southern hybridization with the transposon DNA as a probe.

Assay of GUS activity.

Overnight cultures in LB were transferred to fresh LB, and incubated further. For incubation in IM, log-phase cultures in LB were centrifuged, and cells were washed with IM, before they are resuspended in IM to $OD_{620} \approx 0.5$. The cultures in IM were incubated for an additional 5 to 6 h at 24°C before assay of GUS activity. GUS activity was monitored fluorimetrically as described by Jefferson et al. (1987). Forty-five microliters of the log-phase culture in LB or the induced culture from IM was mixed with an equal volume of 2x assay buffer. After incubation at 37°C for 10 h, GUS activity was measured as described previously (Wei et al. 1992). The background fluorescence of Ea321-G77 (hrcV::Tn5-gusAI) (Wei et al. 1992), which has a gusAI insertion in the opposite direction of hrcV transcription, was subtracted from the readings of hrp::gusAl fusion strains. The corrected fluorescence readings were converted to picounits of GUS activity per CFU. The GUS activity of hrp::Tn5-gusA1 fusions also were determined in tobacco leaf tissues as described previously (Wei et al. 1992).

Plant assays.

Bacteria were grown in LB and harvested at mid-exponential phase. Cells were resuspended in 5 mM potassium phosphate buffer, pH 6.5, harvested again, resuspended in the potassium phosphate buffer to approximately 2 x 108 CFU per ml, unless otherwise specified, and used for HR and pathogenicity assays. Tobacco plants (Nicotiana tabacum L. 'Xanthi') were grown in greenhouse soil mix to a height of 0.9 to 1 m. Bacterial suspensions were infiltrated into each leaf panel of tobacco leaves with needleless hypodermic syringes. The development of the HR was scored after incubation at room temperature for 18 to 36 h. Pathogenicity tests on immature pear fruits were carried out as previously described (Bauer and Beer 1991; Steinberger and Beer 1988).

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NOTE ADDED IN PROOF

A recent BLAST survey of finished and unfinished microbial genomes (available on-line from the NCBI web site) suggests the presence in Pseudomonas aeruginosa PAO1 of a two-component system that is highly similar to the HrpX/HrpY system (31% identity over 474 amino acids for HrpX and 48% identity over 208 amino acids for HrpY)...A related set of proteins exist in the Pseudomonas putida KT2440 genome.

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Bacterial home goal by harpins

Ulla Bonas

ost-pathogen interactions are dynamic and multifactorial; whether a microorganism succeeds or fails in colonizing a potential host depends on factors from both organisms. A successful pathogen has to overcome the defenses of the host. In bacteria that are pathogenic for animals or for plants, particularly Gram-negative organisms, a large number of genes are essential to infect host tissue and establish disease. Expression of these genes is generally controlled by environmental conditions such as temperature, pH, salt concentration and nutrient availability1,2.

Pathogenicity, hypersensitive reaction and elicitors

In the Gram-negative plant pathogens Erwinia, Pseudomonas and Xanthomonas, genes organized in clusters of 25-40 kb are fundamentally involved in any obvious interaction with a plant (for a review see Ref. 3). These genes have been designated hrp (hypersensitive reaction and pathogenicity) because they are essential not only for pathogenicity towards a susceptible host plant, but also for interaction with resistant host varieties and with plants that are not a host for that pathogen. In plants, the hypersensitive reaction (HR) (Ref. 4) is a rapid defense reaction involving localized plant cell death and production of substances such as phenolics and phytoalexins at the site of infection. The HR prevents pathogen spread and thus halts disease development.

In the wild, plants are resistant to the majority of pathogens. The HR, therefore, is an important defense mechanism against all kinds of possible disease agents (bacteria, fungi, nematodes and viruses). It is not only important to interactions of pathogens with nonhost plants, but also to interactions between plants that carry resistance genes and microorganisms that are pathogens for that species.

Although the genes involved in plant defense^{5,6} are becoming better understood, very little is known about the nature of the initial signals and their perception. Induction of the HR in a bacterium-plant interaction requires functional hrp genes and appears to be mediated by signal molecules or 'elicitors'. Recent DNA sequence analyses indicate that several putative Hrp proteins from different species are related and may be involved in a secretion system reminiscent of secretion of Yops (Yersinia outer proteins) in Yersinia⁷⁻¹¹. So far, only one specific elicitor of the HR in a bacterium-plant interaction has been described. The avrD gene from Pseudomonas syringae pv. tomato mediates production of a lowmolecular-mass compound that specifically induces the HR only in the soybean plant (a nonhost) when it carries the corresponding Rpg4 resistance gene5,12.

Harpins

Recently, two bacterial HR-inducing proteins, called 'harpins', were identified in Erwinia amylovora13 and P. syringae pv. syringae14. Although the harpins differ in primary sequence, they have several features in common: they are glycine rich and heat stable, and they both induce an HR in tobacco, a nonhost plant for these bacteria. The genes encoding harpins are localized within the hrp clusters and obviously have a dual role in that they are also required for pathogenicity towards the normal host plant. Both hrp clusters allow nonpathogenic bacteria, such as Escherichia coli, to induce an HR in tobacco after recombinant expression, suggesting that the genes for the tobacco HR elicitors are present within the clusters15,16.

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The first harpin to be identified, harpines, is a cell-envelopeassociated protein encoded by the hrpN gene of Er. amylovora, a pathogen of pear and apple¹³. Recently, He and co-workers¹⁴ have used an elegant approach to identify harpin_{Pss}, which is encoded by the hrpZ gene in the bean pathogen P. s. pv. syringae. Lysates of an expression library in E. coli, made using the cloned P. s. pv. syringae hrp cluster, were directly screened for HR-inducing activity on tobacco leaves. Two proteins were identified, one of which was an amino-terminal deletion of harping with even higher activity than the full-size protein; whether processing occurs during natural infection is not clear. Interestingly, the carboxyl terminus contains two short, direct repeats that are essential for elicitor activity. The activity is in the same range as that of the Erwinia harping; however, to elicit an HR in other plants requires higher levels of the elicitor. He et al. show convincingly that the secretion of harpings by P. s. pv. syringae depends on a product called HrpH that is closely related to proteins in other plant pathogens, and also in animal pathogens such as Yersinia and Shigella, where they are essential for pro-tein secretion^{9,10,14}.

These exciting findings help verify the model that Hrp proteins are involved in the transport of elicitors and virulence factors7. Not surprisingly, the results presented by He and co-workers14 also stimulate many questions. It needs to be shown that harpings is actually secreted when the bacterium interacts with tobacco tissue (the hrp genes were induced in vitro). The concentration needed for HR induction (more than 600 nM) is much higher than one would expect for specific signal molecules. Are harpins toxins? Most importantly, what is their function in pathogenicity, and why do they not elicit an HR in the host plant? Are harpins the only elicitors of nonhost HR in tobacco and possibly in other plants? Is the same mechanism used in tobacco to recognize both the Erwinia and the P. s. pv. syringae harpins? Is host resistance different in mechanism from nonhost resistance? Answers to this fascinating puzzle require the identification of more HR elicitors and their putative plant receptors.

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Initiation and spread of α-herpesvirus infections

Thomas C. Mettenleiter

erpesviruses are large animal viruses with a DNA Lgenome varying from approximately 120 to 250kb. Based on their biological properties, the Herpesviridae have been divided into three subfamilies, the α-, βand y-herpesvirinae, prototypes of which are the human pathogens herpes simplex virus (HSV), cytomegalovirus (HCMV) and Epstein-Barr virus (EBV), respectively. As enveloped viruses, they depend on two consecutive processes for infectious entry into target cells: (1) attachment of free virions to cells and (2) penetration, that is, fusion of virion envelope and cellular cytoplasmic membrane leading to release of the nucleocapsid into the cell. Virion glycoproteins play envelope important roles in both these processes (see Refs 1,2 for recent reviews).

After infection of primary target cells, virus spread can occur by several different mechanisms. Infected cells may release infectious virions that reinitiate infection from outside. In addition, direct viral cell-to-cell spread from primary infected cells to adjacent noninfected cells may occur. In the host, virus may be disseminated by circulating infected cells that adhere to noninfected tissues and transmit infectivity directly. Recent results on HSV and pseudorabies virus (PrV) shed more light on these processes in a-herpesviruses. PrV causes Aujeszky's disease in swine, which is characterized by nervous and respiratory symptoms, and reproductive failure. Unlike HSV, PrV is not pathogenic for humans. However, the two viruses have several features in common, including a broad host range in vitro, and several species besides the known PrV glycoproteins are

the natural host can be infected experimentally. In addition, all of

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related to homologous glycoproteins in HSV (Ref. 1)*.

Attachment

Binding of free infectious virus to target cells involves interactions between virion envelope glycoproteins and cellular virus receptors. Herpes virions contain a large number of different virus-encoded envelope glycoproteins that might participate in attachment. A wellknown example of a cellular herpesvirus receptor is the B-cell membrane protein CR2 (CD21), which binds EBV (Ref. 3). Recent studies have demonstrated that several α- (reviewed in Ref. 1), β- and γherpesviruses4.5 bind to their target cells by interaction of virion components with cell-surface glycosaminoglycans, principally heparan sulfate (HS)6.

^{*}At the 18th International Herpesvirus Workshop, a common nomenclature for α-herpesvirus glycoproteins was agreed on, based on designations of HSV glycoproteins. This nomenclature is used here.

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(54) Title: HYPERSENSITIVE RESPONSE ELICITING DOMAINS AND USE THEREOF

(57) Abstract: The present invention is directed to the structure of an isolated protein or polypeptide which elicits a hypersensitive response in plants as well as an isolated nucleic acid molecule which encodes the hypersensitive response eliciting protein or polypetide. This protein or polypeptide has an acid portion linked to an alpha helix or a pair of spaced apart domains comprising an acidic portion linked to an alpha-helix. This isolated protein or polypeptide and the isolated nucleic acid molecule can be used to impart disease resistance to plants, to enhance plant growth, to control insects, and/or to impart stress resistance to plants. This can be achieved by applying the hypersensitive response elicitor protein or polypeptide in a non-infectious form to plants or plant seeds under conditions effective to impart disease resistance, to enhance plant growth, to control insects, and/or to impart stress resistance to plants or plants grown from the plant seeds. Alternatively, transgenic plants or plant seeds transformed with a nucleic acid molecule encoding a hypersensitive response elicitor protein or polypeptide can be provided and the transgenic plants or plants resulting from the transgenic plant seeds are grown under conditions effective to impart disease resistance, to enhance plant growth, to control insects, and/or to impart stress resistance to plants or plants grown from the plant seeds.

HYPERSENSITIVE RESPONSE ELICITING DOMAINS AND USE THEREOF

This application claims benefit of U.S. Provisional Patent Application Serial No. 60/212,211, filed on June 16, 2000.

FIELD OF THE INVENTION

The present invention relates to hypersensitive response elicitors and their structure. 10

BACKGROUND OF THE INVENTION

Interactions between bacterial pathogens and their plant hosts generally fall into two categories: (1) compatible (pathogen-host), leading to intercellular 15 bacterial growth, symptom development, and disease development in the host plant; and (2) incompatible (pathogen-nonhost), resulting in the hypersensitive response, a particular type of incompatible interaction occurring, without progressive disease symptoms. During compatible interactions on host plants, bacterial populations increase dramatically and progressive symptoms occur. During incompatible 20 interactions, bacterial populations do not increase, and progressive symptoms do not occur.

The hypersensitive response is a rapid, localized necrosis that is associated with the active defense of plants against many pathogens (Kiraly, Z., "Defenses Triggered by the Invader: Hypersensitivity," pages 201-224 in: Plant Disease: An Advanced Treatise, Vol. 5, J.G. Horsfall and E.B. Cowling, ed. Academic Press New York (1980); Klement, Z., "Hypersensitivity," pages 149-177 in: Phytopathogenic Prokaryotes, Vol. 2, M.S. Mount and G.H. Lacy, ed. Academic Press, New York (1982)). The hypersensitive response elicited by bacteria is readily observed as a tissue collapse if high concentrations ($\geq 10^7$ cells/ml) of a limited 30 host-range pathogen like Pseudomonas syringae or Erwinia amylovora are infiltrated into the leaves of nonhost plants (necrosis occurs only in isolated plant cells at lower levels of inoculum) (Klement, Z., "Rapid Detection of Pathogenicity of Phytopathogenic Pseudomonads," Nature 199:299-300; Klement, et al.,

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"Hypersensitive Reaction Induced by Phytopathogenic Bacteria in the Tobacco Leaf," Phytopathology 54:474-477 (1963); Turner, et al., "The Quantitative Relation Between Plant and Bacterial Cells Involved in the Hypersensitive Reaction," Phytopathology 64:885-890 (1974); Klement, Z., "Hypersensitivity," pages 149-177 in Phytopathogenic Prokaryotes, Vol. 2., M.S. Mount and G.H. Lacy, ed. Academic Press, New York (1982)). The capacities to elicit the hypersensitive response in a nonhost and be pathogenic in a host appear linked. As noted by Klement, Z., "Hypersensitivity," pages 149-177 in Phytopathogenic Prokaryotes, Vol. 2., M.S. Mount and G.H. Lacy, ed. Academic Press, New York, these pathogens also cause physiologically similar, albeit delayed, necroses in their interactions with compatible hosts. Furthermore, the ability to produce the hypersensitive response or pathogenesis is dependent on a common set of genes, denoted hrp (Lindgren, P.B., et al., "Gene Cluster of Pseudomonas syringae pv. 'phaseolicola' Controls Pathogenicity of Bean Plants and Hypersensitivity on Nonhost Plants," J. Bacteriol. 168:512-22 (1986); Willis, D.K., et al., "hrp Genes of Phytopathogenic Bacteria," Mol. Plant-Microbe 15 Interact. 4:132-138 (1991)). Consequently, the hypersensitive response may hold clues to both the nature of plant defense and the basis for bacterial pathogenicity.

The hrp genes are widespread in gram-negative plant pathogens, where they are clustered, conserved, and in some cases interchangeable (Willis, D.K., et al., "hrp Genes of Phytopathogenic Bacteria," Mol. Plant-Microbe Interact. 4:132-138 (1991); Bonas, U., "hrp Genes of Phytopathogenic Bacteria," pages 79-98 in: Current Topics in Microbiology and Immunology: Bacterial Pathogenesis of Plants and Animals - Molecular and Cellular Mechanisms, J.L. Dangl, ed. Springer-Verlag, Berlin (1994)). Several hrp genes encode components of a protein secretion pathway similar to one used by Yersinia, Shigella, and Salmonella spp. to secrete proteins essential in animal diseases (Van Gijsegem, et al., "Evolutionary Conservation of Pathogenicity Determinants Among Plant and Animal Pathogenic Bacteria," Trends Microbiol. 1:175-180 (1993)). In E. amylovora, P. syringae, and P. solanacearum, hrp genes have been shown to control the production and secretion of glycine-rich, protein elicitors of the hypersensitive response (He, S.Y., et al. "Pseudomonas 30 Syringae pv. Syringae HarpinPss: a Protein that is Secreted via the Hrp Pathway and Elicits the Hypersensitive Response in Plants," Cell 73:1255-1266 (1993), Wei, Z.-H.,

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et al., "HrpI of Erwinia amylovora Functions in Secretion of Harpin and is a Member of a New Protein Family," <u>J. Bacteriol.</u> 175:7958-7967 (1993); Arlat, M. et al. "PopA1, a Protein Which Induces a Hypersensitive-like Response on Specific Petunia Genotypes, is Secreted via the Hrp Pathway of Pseudomonas solanacearum," <u>EMBO</u> J. 13:543-553 (1994)).

The first of these proteins was discovered in E. amylovora Ea321, a bacterium that causes fire blight of rosaceous plants, and was designated harpin (Wei, Z.-M., et al, "Harpin, Elicitor of the Hypersensitive Response Produced by the Plant Pathogen Erwinia amylovora," Science 257:85-88 (1992)). Mutations in the encoding hrpN gene revealed that harpin is required for E. amylovora to elicit a hypersensitive response in nonhost tobacco leaves and incite disease symptoms in highly susceptible pear fruit. The P. solanacearum GMI1000 PopA1 protein has similar physical properties and also elicits the hypersensitive response in leaves of tobacco, which is not a host of that strain (Arlat, et al. "PopA1, a Protein Which Induces a Hypersensitive-like Response on Specific Petunia Genotypes, is Secreted via the Hrp Pathway of Pseudomonas solanacearum," EMBO J. 13:543-53 (1994)). However, P. solanacearum popA mutants still elicit the hypersensitive response in tobacco and incite disease in tomato. Thus, the role of these glycine-rich hypersensitive response elicitors can vary widely among gram-negative plant pathogens.

Other plant pathogenic hypersensitive response elicitors have been isolated, cloned, and sequenced. These include: Erwinia chrysanthemi (Bauer, et. al., "Erwinia chrysanthemi Harpingeh: Soft-Rot Pathogenesis," MPMI 8(4): 484-91 (1995)); Erwinia carotovora (Cui, et. al., "The RsmA" Mutants of Erwinia carotovora subsp. carotovora Strain Ecc71 Overexpress hrpNge and Elicit a Hypersensitive Reaction-like Response in Tobacco Leaves," MPMI 9(7): 565-73 (1966)); Erwinia stewartii (Ahmad, et. al., "Harpin is not Necessary for the Pathogenicity of Erwinia stewartii on Maize," 8th Int'l. Cong. Molec. Plant-Microb, Inter. July 14-19, 1996 and Ahmad, et. al., "Harpin is not Necessary for the Pathogenicity of Erwinia stewartii on Maize," Ann. Mtg. Am. Phytopath. Soc. July 27-31, 1996); and Pseudomonas syringae pv. syringae (WO 94/26782 to Cornell Research Foundation, Inc.).

The present invention is a further advance in the effort to identify and characterize hypersensitive response elicitor proteins.

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SUMMARY OF THE INVENTION

One aspect of the present invention is directed to an isolated hypersensitive response elicitor protein comprising a pair of spaced apart domains, with each comprising an acid portion linked to an alpha-helix.

Another embodiment of the present invention relates to an isolated hypersensitive response elicitor protein comprising an acid portion linked to an alphahelix.

Nucleic acid molecules encoding either of these proteins as well as vectors, bost cells, transgenic plants, and transgenic plant seeds containing those nucleic acid molecules are also disclosed.

The protein of the present invention can be used to impart disease resistance to plants, to enhance plant growth, to control insects, and/or impart stress resistance. This involves applying the protein to plants or plant seeds under conditions effective to impart disease resistance, to enhance plant growth, to control insects, and/or impart stress resistance to plants or plants grown from the plant seeds.

As an alternative to applying the protein to plants or plant seeds in order to impart disease resistance, to enhance plant growth, to control insects on plants, and/or impart stress resistance, transgenic plants or plant seeds can be utilized. When utilizing transgenic plants, this involves providing a transgenic plant transformed with a nucleic acid molecule encoding the protein of the present invention and growing the plant under conditions effective to impart disease resistance, to enhance plant growth, to control insects, and/or to impart stress resistance to the plants or plants grown from the plant seeds. Alternatively, a transgenic plant seed transformed with the nucleic acid molecule encoding the protein of the present invention can be provided and planted in soil. A plant is then propagated under conditions effective to impart disease resistance, to enhance plant growth, to control insects, and/or to impart stress resistance to plants or plants grown from the plant seeds.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a schematic drawing showing the construction of a universal expression cassette for a hypersensitive response domain.

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DETAILED DESCRIPTION OF THE INVENTION

The present invention is directed to an isolated hypersensitive response elicitor protein comprising a pair of spaced apart domains, with each comprising an acid portion linked to an alpha-helix. The acidic portion is a polypeptide with 10 or more amino acids, is rich in acidic amino acids, and has a pI below 5.0. The acidic portion has a secondary structure in the form of a beta-sheet or a beta-turn. The secondary structure of this unit can also be in an unordered form.

The alpha-helix portion of the present invention is a polypeptide with 10 or more amino acids. Its secondary structure is in the form of a stable alpha-helix.

Another embodiment of the present invention relates to an isolated hypersensitive response elicitor protein comprising an acid portion linked to an alphahelix.

Both of these proteins are capable of eliciting a hypersensitive response.

The alpha helix is a common structural motif of proteins in which a linear sequence of amino acid folds into a right-handed helix stabilized by internal hydrogen bonding between backbone atoms.

The acidic motif includes a certain combination of amino acids in which a linear sequence with a pI below 5.0 folds into a β sheet, coil, or thin structures but not an alpha helix of secondary structure.

The hypersensitive response elicitor polypeptides or proteins according to the present invention can be derived from hypersensitive response elicitor polypeptides or proteins of a wide variety of fungal and bacterial pathogens. Such polypeptides or proteins are able to elicit local necrosis in plant tissue contacted by the elicitor. Examples of suitable bacterial sources of polypeptide or protein elicitors

include Erwinia, Pseudomonas, and Xanthamonas species (e.g., the following bacteria: Erwinia amylovora, Erwinia chrysanthemi, Erwinia stewartii, Erwinia carotovora, Pseudomonas syringae, Pseudomonas solancearum, Xanthomonas campestris, and mixtures thereof). In addition to hypersensitive response elicitors from these Gram negative bacteria, it is possible to use elicitors from Gram positive bacteria. One example is Clavibacter michiganensis subsp. sepedonicus.

An example of a fungal source of a hypersensitive response elicitor protein or polypeptide is Phytophthora. Suitable species of Phytophthora include Phytophthora parasitica, Phytophthora cryptogea, Phytophthora cinnamomi, Phytophthora capsici, Phytophthora megasperma, and Phytophthora citrophthora.

The hypersensitive response elicitor polypeptide or protein from Erwinia chrysanthemi has an amino acid sequence corresponding to SEQ. ID. No. 1 as follows:

Met Gln Ile Thr Ile Lys Ala His Ile Gly Gly Asp Leu Gly Val Ser 1 5 10 15 15 Gly Leu Gly Ala Gln Gly Leu Lys Gly Leu Asn Ser Ala Ala Ser Ser 20 25 30 Leu Gly Ser Ser Val Asp Lys Leu Ser Ser Thr Ile Asp Lys Leu Thr 20 Ser Ala Leu Thr Ser Met Met Phe Gly Gly Ala Leu Ala Gln Gly Leu Gly Ala Ser Ser Lys Gly Leu Gly Met Ser Asn Gln Leu Gly Gln Ser 65 70 75 80 Phe Gly Asn Gly Ala Gln Gly Ala Ser Asn Leu Leu Ser Val Pro Lys 25 Ser Gly Gly Asp Ala Leu Ser Lys Met Phe Asp Lys Ala Leu Asp Asp Leu Leu Gly His Asp Thr Val Thr Lys Leu Thr Asn Gln Ser Asn Gln 30 Leu Ala Asn Ser Met Leu Asn Ala Ser Gln Met Thr Gln Gly Asn Met Asn Ala Phe Gly Ser Gly Val Asn Asn Ala Leu Ser Ser Ile Leu Gly Asn Gly Leu Gly Gln Ser Met Ser Gly Phe Ser Gln Pro Ser Leu Gly 35

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	Ala	Gly	Gly	Leu 180	Gln	Gly	Leu	Ser	Gly 185	Ala	Gly	Ala	Phe	Asn 190	Gln	Leu
	Gly	Asn	Ala 195	Ile	Gly	Met	Gly	Val 200	Gly	Gln	Asn	Ala	Ala 205	Leu	Ser	Ala
5	Leu	Ser 210	Asn	Val	Ser	Thr	His 215	Val	Ąsp	Gly	Asn	Asn 220	Arg	His	Phe	Val
	Авр 225	Lys	Glu	Ąsp	Arg	Gly 230	Met	Ala	Lys	Glu	Ile 235	Gly	Gln	Phe	Met	Asp 240
10	Gln	Tyr	Pro	Glu	Ile 245	Phe	Gly	Lys	Pro	Glu 250	Tyr	Gln	Lys	Авр	Gly 255	Trp
.*	Ser	Ser	Pro	Lув 260	Thr	Авр	Авр	Lys	Ser 265	Trp	Ala	Lys	Ala	Leu 270	Ser	ГЛЯ
	Pro	Asp	Asp 275		Gly	Met	Thr	Gly 280	Ala	ser	Met	Asp	Lys 285	Phe	Arg	gln
15	Ala	Met 290		Met	Ile	Lys	Ser 295	Als	Val	Ala	Gly	300 300	Th:	c Gly	ABI	1 Thr
	Asn 305		Asn	Leu	Arg	Gly 310	r Ala	Gly	, Gl	y Ale	315	Let	ı Gl	y Ile	а Авј	9 Ala 320
20	Ala	Val	. Val	. Gly	Asj 32	Lys S	Ile	. Ala	a Ası	n Mei 33	sei O	r Le	u Gl	у Гу	8 Le	u Ala 5
	Ası	Ala														

This hypersensitive response elicitor polypeptide or protein has a molecular weight of 34 kDa, is heat stable, has a glycine content of greater than 16%, and contains substantially no cysteine. The *Erwinia chrysanthemi* hypersensitive response elicitor polypeptide or protein is encoded by a DNA molecule having a nucleotide sequence corresponding to SEQ. ID. No. 2 as follows:

GCGTTTATCC CGGGTGAACG TGCTATGACC GACAGCATCA CGGTATTCGA CACCGTTACG 60
GCGTTTATGG CCGCGATGAA CCGGCATCAG GCGGCGCGCT GGTCGCCGCA ATCCGGCGTC 120
GATCTGGTAT TTCAGTTTGG GGACACCGGG CGTGAACTCA TGATGCAGAT TCAGCCGGGG 180
CAGCAATATC CCGGCATGTT GCGCACGCTG CTCGCTCGTC GTTATCAGCA GGCGGCAGAG 240
TGCGATGGCT GCCATCTGTG CCTGAACGGC AGCGATGTAT TGATCCTCTG GTGGCCGCTG 300
ACGTTGCGATC CCGGCAGTTA TCCGCAGGTG ATCGAACGTT TGTTTGAACT GGCGGGAATG 360
ACGTTGCCGT CGCTATCCAT AGCACCGACG GCGCGTCCGC AGACAGGGAA CGGACGCGCC 420
CGATCATTAA GATAAAGGCG GCTTTTTTTA TTGCAAAACG GTAACGGTGA GGAACCGTTT 460

	CACCGTCGGC GTCACTCAGT AACAAGTATC CATCATGATG CCTACATCGG GATCGGCGTG	540
		600
	GGCATCCGTT GCAGATACTT TTGCGAACAC CTGACATGAA TGAGGAAACG AAATTATGCA	800
	ANTIACGATC ARAGCGCACA TCGGCGGTGA TTTGGGCGTC TCCGGTCTGG GGCTGGGTGC	660
	TCAGGGACTG AAAGGACTGA ATTCCGCGGC TTCATCGCTG GGTTCCAGCG TGGATAAACT	720
5	GAGCAGCACC ATCGATAAGT TGACCTCCGC GCTGACTTCG ATGATGTTTG GCGGCGCGCT	780
	GGCGCAGGGG CTGGGCGCCA GCTCGAAGGG GCTGGGGATG AGCAATCAAC TGGGCCAGTC	840
	TITCOGCAAT GGCGCGCAGG GTGCGAGCAA CCTGCTATCC GTACCGAAAT CCGGCGGCGA	900
	TGCGTTGTCA AAAATGTTTG ATAAAGCGCT GGACGATCTG CTGGGTCATG ACACCGTGAC	960
	CAAGCTGACT AACCAGAGCA ACCAACTGGC TAATTCAATG CTGAACGCCA GCCAGATGAC	1020
10	CCAGGGTAAT ATGAATGCGT TCGGCAGCGG TGTGAACAAC GCACTGTCGT CCATTCTCGG	1080
	CAACGGTCTC GGCCAGTCGA TGAGTGGCTT CTCTCAGCCT TCTCTGGGGG CAGGCGGCTT	1140
	GCAGGCCTG AGCGGCGCGG GTGCATTCAA CCAGTTGGGT AATGCCATCG GCATGGGCGT	1200
	GGGGCAGAAT GCTGCGCTGA GTGCGTTGAG TAACGTCAGC ACCCACGTAG ACGGTAACAA	1260
	COGCCACTIT GTAGATAAAG AAGATCGCGG CATGGCGAAA GAGATCGGCC AGTTTATGGA	1320
15	TCAGTATCCG GAAATATTCG GTAAACCGGA ATACCAGAAA GATGGCTGGA GTTCGCCGAA	1380
	GACOGACGAC AAATCCTGGG CTAAAGCGCT GAGTAAACCG GATGATGACG GTATGACCGG	1440
	COCCAGCATG GACAAATTCC GTCAGGCGAT GGGTATGATC AAAAGCGCGG TGGCGGGTGA	1500
	TACCGGCAAT ACCAACCTGA ACCTGCGTGG CGCGGGCGGT GCATCGCTGG GTATCGATGC	1560
	GGCTGTCGTC GGCGATAAAA TAGCCAACAT GTCGCTGGGT AAGCTGGCCA ACGCCTGATA	1620
20	ATCTGTGCTG GCCTGATAAA GCGGAAAGGA AAAAAGAGAC GGGGAAGCCT GTCTCTTTTC	1680
	TIATTATGCG GTTTATGCGG TTACCTGGAC CGGTTAATCA TCGTCATCGA TCTGGTACAA	1740
	ACGCACATTT TCCCGTTCAT TCGCGTCGTT ACGCGCCACA ATCGCGATGG CATCTTCCTC	1800
	GTCGCTCAGA TTGCGCGGCT GATGGGGAAC GCCGGGTGGA ATATAGAGAA ACTCGCCGGC	1860
	CAGATEGAGA CACETCTECE ATARATCTET GCCGTAACET GTTTCTATCC GCCCCTTTAG	1920
25	CAGATAGATT GCGGTTTCGT AATCAACATG GTAATGCGGT TCCGCCTGTG CGCCGGCCGG	1980
	GATCACCACA ATATTCATAG ARAGCTGTCT TGCACCTACC GTATCGCGGG AGATACCGAC	2040
	AAAATAGGGC AGTTTTTGCG TGGTATCCGT GGGGTGTTCC GGCCTGACAA TCTTGAGTTG	210
	GITCGTCATC ATCTTTCTCC ATCTGGGCGA CCTGATCGGT T	214

The hypersensitive response elicitor from Erwinia chrysanthemi has 2 hypersensitive response eliciting domains. The first domain extends, within SEQ.

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ID. No. 1, from amino acid 69 to amino acid 122, particularly from amino acid 85 to amino acid 116. The acidic unit in the first domain extends, within SEQ. ID. No. 1, from amino acid 69 to amino acid 102, particularly from amino acid 85 to amino acid 102. The alpha-helix in the first domain extends, within SEQ. ID. No. 1, from amino acid 102 to amino acid 122, particularly from amino acid 102 to amino acid 116. The second domain extends, within SEQ. ID. No. 1, from amino acid 251 to amino acid 299, particularly from amino acid 256 to amino acid 292. The acidic unit in the second domain extends, within SEQ. ID. No. 1, from amino acid 251 to amino acid 279, particularly from amino acid 261 to amino acid 279. The alpha-helix in the second domain extends, within SEQ. ID. No. 1, from amino acid 279 to amino acid 299, particularly from amino acid 279 to amino acid 279 to amino acid 299, particularly from amino acid 279 to amino acid 292.

The hypersensitive response elicitor polypeptide or protein derived from Erwinia amylovora has an amino acid sequence corresponding to SEQ. ID. No. 3 as follows:

Met Ser Leu Asn Thr Ser Gly Leu Gly Ala Ser Thr Met Gln Ile Ser Ile Gly Gly Ala Gly Gly Asn Asn Gly Leu Leu Gly Thr Ser Arg Gln 20 25 30 20 Asn Ala Gly Leu Gly Gly Asn Ser Ala Leu Gly Leu Gly Gly Gly Asn 35 40 45 Gln Asn Asp Thr Val Asn Gln Leu Ala Gly Leu Leu Thr Gly Met Met Met Met Met Ser Met Met Gly Gly Gly Leu Met Gly Gly Leu 25 Gly Gly Leu Gly Asn Gly Leu Gly Gly Ser Gly Gly Leu Gly Glu Gly Leu Ser Asn Ala Leu Asn Asp Met Leu Gly Gly Ser Leu Asn Thr 30 Leu Gly Ser Lys Gly Gly Asn Asn Thr Thr Ser Thr Thr Asn Ser Pro Leu Asp Gln Ala Leu Gly Ile Asn Ser Thr Ser Gln Asn Asp Asp Ser Thr Ser Gly Thr Asp Ser Thr Ser Asp Ser Ser Asp Pro Met Gln Gln 35 160 150

							_	10									
<i>:</i> *	Leu	Leu,	Lys	Met	Phe 165	Ser	Glu i	Ile !	Met	Gln 170	Ser	Leu	Phe	Gly	Авр 175	Gl	У
	Gln	qaA	Gly	Thr 180	Gln	Gĵ	Ser	Ser	Ser 185	Gly	Gly	Lys	Gln	Pro 190	Thr	G)	.u
5	Gly	Glu	Gln 195	Asn	Ala	Tyr	ГЛВ	1ув 200	Gly	Val	Thr	qaA	Ala 205	Leu	Sex	: G]	ГУ
		210					Ser 215					220					
10	225					230	Ala				230	,				_	••
					245		Asn			250	,					_	
				260			Thr		265					2,	•		
15			275	i			Thr	280					20	_			
		290					Ala 295					30	U				
20	305					310					31						
					325	5	- Asi			33	0				_	-55	
				34	D		y Met		34	5				3	30		
25			35	5			e Ly	36	U				3	0.5			
		370	0				37	5					6 U				Asp
30	Ala 385		t Me	t Al	a Gl	у Ав 39	p Al O	a Il	e As	en A	M na E	et A 95	la I	eu (3ly	ГÀВ	Leu 400
	Gly	/ Al	a Al	a						•							

This hypersensitive response elicitor polypeptide or protein has a molecular weight of about 39 kDa, has a pI of approximately 4.3, and is heat stable at 100°C for at least 10 minutes. This hypersensitive response elicitor polypeptide or protein has substantially no cysteine. The hypersensitive response elicitor polypeptide or protein derived from Erwinia amylovora is more fully described in Wei, Z.-M., R. J. Laby, C. H. Zumoff,

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D. W. Bauer, S.-Y. He, A. Collmer, and S. V. Beer, "Harpin, Elicitor of the Hypersensitive Response Produced by the Plant Pathogen Erwinia amylovora," Science 257:85-88 (1992), which is hereby incorporated by reference. The DNA molecule encoding this polypeptide or protein has a nucleotide sequence corresponding to SEQ. ID. No. 4 as follows:

AAGCTTCGGC ATGGCACGTT TGACCGTTGG GTCGGCAGGG TACGTTTGAA TTATTCATAA 60 GAGGAATACG TTATGAGTCT GAATACAAGT GGGCTGGGAG CGTCAACGAT GCAAATTTCT 120 ATCGGCGGTG CGGGCGGAAA TAACGGGTTG CTGGGTACCA GTCGCCAGAA TGCTGGGTTG 180 GGTGGCAATT CTGCACTGGG GCTGGGCGGC GGTAATCAAA ATGATACCGT CAATCAGCTG 240 GCTGGCTTAC TCACCGGCAT GATGATGATG ATGAGCATGA TGGGCGGTGG TGGGCTGATG 300 GGCGGTGGCT TAGGCGGTGG CTTAGGTAAT GGCTTGGGTG GCTCAGGTGG CCTGGGCGAA 360 GGACTGTCGA ACGCGCTGAA CGATATGTTA GGCGGTTCGC TGAACACGCT GGGCTCGAAA 420 GGCGGCAACA ATACCACTTC AACAACAAAT TCCCCGCTGG ACCAGGCGCT GGGTATTAAC 480 TCAACGTCCC AAAACGACGA TTCCACCTCC GGCACAGATT CCACCTCAGA CTCCAGCGAC 540 CCGATGCAGC AGCTGCTGAA GATGTTCAGC GAGATAATGC AAAGCCTGTT TGGTGATGGG 600 CAAGATGGCA CCCAGGGCAG TTCCTCTGGG GGCAAGCAGC CGACCGAAGG CGAGCAGAAC 660 GCCTATAAAA AAGGAGTCAC TGATGCGCTG TCGGGCCTGA TGGGTAATGG TCTGAGCCAG 720 CTCCTTGGCA ACGGGGGACT GGGAGGTGGT CAGGGCGGTA ATGCTGGCAC GGGTCTTGAC 780 GGTTCGTCGC TGGGCGGCAA AGGGCTGCAA AACCTGAGCG GGCCGGTGGA CTACCAGCAG 840 TTAGGTAACG CCGTGGGTAC CGGTATCGGT ATGAAAGCGG GCATTCAGGC GCTGAATGAT 900 ATCGGTACGC ACAGGCACAG TTCAACCCGT TCTTTCGTCA ATAAAGGCGA TCGGGCGATG 960 GCGAAGGAAA TCGGTCAGTT CATGGACCAG TATCCTGAGG TGTTTGGCAA GCCGCAGTAC 1020 CAGAAAGGCC CGGGTCAGGA GGTGAAAACC GATGACAAAT CATGGGCAAA AGCACTGAGC 1080 AAGCCAGATG ACGACGGAAT GACACCAGCC AGTATGGAGC AGTTCAACAA AGCCAAGGGC 1140 ATGATCAAAA GGCCCATGGC GGGTGATACC GGCAACGGCA ACCTGCAGGC ACGCGGTGCC 1200 GGTGGTTCTT CGCTGGGTAT TGATGCCATG ATGCCCGGTG ATGCCCATTAA CAATATGGCA 1260 1288 CTTGGCAAGC TGGGCGCGGC TTAAGCTT

The hypersensitive response elicitor from Erwinia amylovora has 2 hypersensitive response eliciting domains. The first domain extends, within SEQ. ID. No. 3, from amino acid 32 to amino acid 74, particularly from amino acid 45 to amino

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acid 68. The acidic unit in the first domain extends, within SEQ. ID. No. 3, from amino acid 32 to amino acid 57, particularly from amino acid 45 to amino acid 57. The alpha-helix in the first domain extends, within SEQ. ID. No. 3, from amino acid 57 to amino acid 74, particularly from amino acid 57 to amino acid 68. The second domain extends, within SEQ. ID. No. 3, from amino acid 130 to amino acid 180, particularly from amino acid 145 to amino acid 170. The acidic unit in the second domain extends, within SEQ. ID. No. 3, from amino acid 130 to amino acid 157, particularly from amino acid 145 to amino acid 157. The alpha-helix in the second domain extends, within SEQ. ID. No. 3, from amino acid 157 to amino acid 180, particularly from amino acid 157 to amino acid 170.

Another potentially suitable hypersensitive response elicitor from Erwinia amylovora is disclosed in U.S. Patent Application Serial No. 09/120,927, which is hereby incorporated by reference. The protein is encoded by a DNA molecule having a nucleic acid sequence of SEQ. ID. No. 5 as follows:

15 ATGTCAATTC TTACGCTTAA CAACAATACC TCGTCCTCGC CGGGTCTGTT CCAGTCCGGG 60 GGGGACAACG GGCTTGGTGG TCATAATGCA AATTCTGCGT TGGGGCAACA ACCCATCGAT 120 CGGCARACCA TTGAGCARAT GGCTCARTTA TTGGCGGAAC TGTTARAGTC ACTGCTATCG 180 20 CCACAATCAG GTAATGCGGC AACCGGAGCC GGTGGCAATG ACCAGACTAC AGGAGTTGGT 240 NACGCTGGCG GCCTGAACGG ACGANAAGGC ACAGCAGGAA CCACTCCGCA GTCTGACAGT 300 25 CAGAACATGC TGAGTGAGAT GGGCAACAAC GGGCTGGATC AGGCCATCAC GCCCGATGGC 360 CAGGGCGGCG GGCAGATCGG CGATAATCCT TTACTGAAAG CCATGCTGAA GCTTATTGCA 420 CGCATGATGG ACGGCCAAAG CGATCAGTTT GGCCAACCTG GTACGGGCAA CAACAGTGCC 480 30 TCTTCCGGTA CTTCTTCATC TGGCGGTTCC CCTTTTAACG ATCTATCAGG GGGGAAGGCC 540 CCTTCCGGCA ACTCCCCTTC CGGCAACTAC TCTCCCGTCA GTACCTTCTC ACCCCCATCC 600 35 ACGCCARCET CCCCTACCTC ACCGCTTGAT TTCCCTTCTT CTCCCACCAA AGCAGCCGGG 660 GGCAGCACGC CGGTAACCGA TCATCCTGAC CCTGTTGGTA GCGCGGGCAT CGGGGCCGGA 720 AATTCGGTGG CCTTCACCAG CGCCGGCGCT AATCAGACGG TGCTGCATGA CACCATTACC 780 40 GTGAAAGCGG GTCAGGTGTT TGATGGCAAA GGACAAACCT TCACCGCCGG TTCAGAATTA 840 GGCGATGGCG GCCAGTCTGA AAACCAGAAA CCGCTGTTTA TACTGGAAGA CGGTGCCAGC 900 45 CTGAAAAACG TCACCATGGG CGACGACGGG GCGGATGGTA TTCATCTTTA CGGTGATGCC 960 AAAATAGACA ATCTGCACGT CACCAACGTG GGTGAGGACG CGATTACCGT TAAGCCAAAC 1020 AGCGCGGGCA AAAAATCCCA CGTTGAAATC ACTAACAGTT CCTTCGAGCA CGCCTCTGAC 1080 50

	ANGATOCTEC AGCTGAATEC CGATACTAAC CTGAGCETTG ACAACETGAA GGCCAAAGAC	1140
	TTTGGTACTT TTGTACGCAC TAACGGCGGT CAACAGGGTA ACTGGGATCT GAATCTGAGC	1200
5	CATATCAGCG CAGAAGACGG TAAGTTCTCG TTCGTTAAAA GCGATAGCGA GGGGCTAAAC	1260
	GTCAATACCA GTGATATCTC ACTGGGTGAT GTTGAAAACC ACTACAAAGT GCCGATGTCC	1320
10	GCCAACCTGA AGGTGGCTGA ATGA	1344
	See GenBank Accession No. U94513. The isolated DNA molecule of the pr	esent
	invention encodes a hypersensitive response elicitor protein or polypeptide h	aving an
		_
	amino acid sequence of SEQ. ID. No. 6 as follows:	
15	Met Ser Ile Leu Thr Leu Asn Asn Asn Thr Ser Ser Ser Pro 1 5 10	13
20	Phe Gln Ser Gly Gly Asp Asn Gly Leu Gly Gly His Asn Ala 20 25 30	
	Ala Leu Gly Gln Gln Pro Ile Asp Arg Gln Thr Ile Glu Gln 35 40 45	
25	Gln Leu Leu Ala Glu Leu Leu Lys Ser Leu Leu Ser Pro Gln 50 55 60	
30	Asn Ala Ala Thr Gly Ala Gly Gly Asn Asp Gln Thr Thr Gly 65 70 75	00
50	Asn Ala Gly Gly Leu Asn Gly Arg Lys Gly Thr Ala Gly Thr 85 90	33
35	Gln Ser Asp Ser Gln Asn Met Leu Ser Glu Met Gly Asn Asn 100 105 110	•
	Asp Gln Ala Ile Thr Pro Asp Gly Gln Gly Gly Gln Ile 115 120 125	
40	130 135 140	•
45	Gly Gln Ser Asp Gln Phe Gly Gln Pro Gly Thr Gly Asn As 145 150 155	
	Ser Ser Gly Thr Ser Ser Ser Gly Gly Ser Pro Phe Ash As 165 170	
50	180	•
•	Val Ser Thr Phe Ser Pro Pro Ser Thr Pro Thr Ser Pro Ti 195 200 205	
55	Leu Asp Phe Pro Ser Ser Pro Thr Lys Ala Ala Gly Gly S 210 215 220	er Thr Pro

· · ·	225			•		230					Ala 235			ı	,	220
5	Asn	Ser	Val	Ala	Phe 245	Thr	Ser	Ala	Gly	Ala 250	Asn	Gln	Thr	Val	Leu 255	His
	Asp	Thr	Ile	Thr 260	Val	Lys	Ala	Gly	Gln 265	Val	Phe	qaA	Gly	Lув 270	Gly	Gln
10	Thr	Phe	Thr 275		Gly	Ser	Glu	Leu 280	Gly	Авр	Gly	Gly	Gln 285	Ser	Glu	Asn
	Gln	Lys 290		Leu	Phe	Ile	Leu 295	Glu	qaA	Gly	alA v	Ser 300	Leu	Lys	Asn	Val
15 .	Thr 305		gly	Asp	Asp	Gly 310	Ala	Авр	Gly	Ile	315	Leu	Тух	Gly	Asp	Ala 320
20	Lys	Ile	Asp	Asn	Leu 325	His	Val	Thr	Ası	1 Val 330	o L Gly	r Glu	l Ası	Ala	335	Thr
	Val	Lув	Pro	340	Ser	Ala	Gly	Lys	Ly:	8e:	r Hie	va.	Gl:	350	Thi	r Asn
25	Ser	Ser	? Phe		His	Ala	Ser	360	Lyı)	s Il	e Let	a Gla	1 Le	u Aas S	Ala	а Азр
	Thr	Э ДВІ 370		ı Sei	· Val	Asr	Э Авг 375	val	L Ly:	s Al	а Ьу	8 As	p Ph O	e Gl	y Th	r Phe
30	Val 385		Thi	C Asi	a Gl	Gly 390		ı Glı	n Gl	у Ав	n Tr 39	p As 5	рЬе	u As	n Le	u Ser 400
35	His	, Ile	e Se	r Ala	40:		p Gl	y Ly	e Pb	e Se 43	r Ph	e Va	l Ly	ra Se	x As	p Ser 15
	Glu	ı Gl	y Le	1 Asi 42		l As	n Th	r Se	r As 42	p II 15	le 86	r Le	eu G	Ly As 43	np Va 30	al Glu
40 .	Ası	n Hi	в Ту 43		e Va	l Pr	o Me	t Se 44	r Al O	ia Ai	en Le	eu Ly	78 V:	al Ai 45	la G	lu

This protein or polypeptide is acidic, rich in glycine and serine, and lacks cysteine. It is also heat stable, protease sensitive, and suppressed by inhibitors of plant metabolism. The protein or polypeptide of the present invention has a predicted molecular size of ca. 4.5 kDa.

This hypersensitive response elicitor from Erwinia amylovora has 2 hypersensitive response eliciting domains. The first domain extends, within SEQ. ID. No. 6, from amino acid 5 to amino acid 64, particularly from amino acid 31 to amino acid 57. The acidic unit in the first domain extends, within SEQ. ID. No. 6, from amino acid 5 to amino acid 45, particularly from amino acid 31 to amino acid 45. The

alpha-helix in the first domain extends, within SEQ. ID. No. 6, from amino acid 45 to amino acid 64, particularly from amino acid 45 to amino acid 64. The second domain extends, within SEQ. ID. No. 6, from amino acid 103 to amino acid 146, particularly from amino acid 116 to amino acid 140. The acidic unit in the second domain extends, within SEQ. ID. No. 6, from amino acid 103 to amino acid 131, particularly from amino acid 116 to amino acid 131. The alpha-helix in the second domain extends, within SEQ. ID. No. 6, from amino acid 131 to amino acid 146, particularly from amino acid 131 to amino acid 131 to amino acid 140.

Another potentially suitable hypersensitive response elicitor from Erwinia amylovora is disclosed in U.S. Patent Application Serial No. 09/120,663, which is hereby incorporated by reference. The protein is encoded by a DNA molecule having a nucleic acid sequence of SEQ. ID. No. 7 as follows:

ATGGAATTAA AATCACTGGG AACTGAACAC AAGGCGGCAG TACACACAGC GGCGCACAAC 60 15 CCTGTGGGGC ATGGTGTTGC CTTACAGCAG GGCAGCAGCA GCAGCAGCCC GCAAAATGCC 120 GCTGCATCAT TGGCGGCAGA AGGCAAAAAT CGTGGGAAAA TGCCGAGAAT TCACCAGCCA 180 TCTACTGCGG CTGATGGTAT CAGCGCTGCT CACCAGCAAA AGAAATCCTT CAGTCTCAGG 240 20 GGCTGTTTGG GGACGAAAAA ATTTTCCAGA TCGGCACCGC AGGGCCAGCC AGGTACCACC 300 CACAGCAAAG GGGCAACATT GCGCGATCTG CTGGCGCGGG ACGACGGCGA AACGCAGCAT 360 25 GAGGCGGCCG CGCCAGATGC GGCGCGTTTG ACCCGTTCGG GCGGCGTCAA ACGCCGCAAT 420 ATGGACGACA TGGCCGGGCG GCCAATGGTG AAAGGTGGCA GCGGCGAAGA TAAGGTACCA 480 ACGCAGCAAA AACGGCATCA GCTGAACAAT TTTGGCCAGA TGCGCCAAAC GATGTTGAGC 540 30 AAAATGGCTC ACCOGCTTC AGCCAACGCC GGCGATCGCC TGCAGCATTC ACCGCCGCAC 600 ATCCCGGGTA GCCACCACGA AATCAAGGAA GAACCGGTTG GCTCCACCAG CAAGGCAACA 660 35 ACGGCCCACG CAGACAGAGT GGAAATCGCT CAGGAAGATG ACGACAGCGA ATTCCAGCAA 720 CTGCATCAAC AGCGGCTGGC GCGCGAACGG GAAAATCCAC CGCAGCCGCC CAAACTCGGC 780 GTTGCCACAC CGATTAGCGC CAGGTTTCAG CCCAAACTGA CTGCGGTTGC GGAAAGCGTC 840 40 CTTGAGGGGA CAGATACCAC GCAGTCACCC CTTAAGCCGC AATCAATGCT GAAAGGAAGT 900 GGAGCCGGGG TAACGCCGCT GGCGGTAACG CTGGATAAAG GCAAGTTGCA GCTGGCACCG 960 45 GATAATCCAC CCGCGCTCAA TACGTTGTTG AAGCAGACAT TGGGTAAAGA CACCCAGCAC 1020 TATCTGGGGC ACCATGCCAG CAGCGACGGT AGCCAGCATC TGCTGCTGGA CAACAAAGGC 1080 CACCTGTTTG ATATCAAAAG CACCGCCACC AGCTATAGCG TGCTGCACAA CAGCCACCCC 1140 GGTGAGATAA AGGGCAAGCT GGCGCAGGCG GGTACTGGCT CCGTCAGCGT AGACGGTAAA 1200

	·	
	AGCGGCAAGA TCTCGCTGGG GAGCGGTACG CAAAGTCACA ACTIO	1260
	CCGGGGGAAG CGCACCGTTC CTTATTAACC GGCATTTGGC AGCATCCTGC TGGCGCAGCG	1320
5	CGGCCGCAGG GCGAGTCAAT CCGCCTGCAT GACGACAAAA TTCATATCCT GCATCCGGAG	1380
	CTGGGCGTAT GGCAATCTGC GGATAAAGAT ACCCACAGCC AGCTGTCTCG CCAGGCAGAC	1440
_	GGTAAGCTCT ATGCGCTGAA AGACAACCGT ACCCTGCAAA ACCTCTCCGA TAATAAATCC	1500
0	TCAGAAAAGC TGGTCGATAA AATCAAATCG TATTCCGTTG ATCAGCGGGG GCAGGTGGCG	1560
	ATCCTGACGG ATACTCCCGG CCGCCATAAG ATGAGTATTA TGCCCTCGCT GGATGCTTCC	1620
.5	CCGGAGAGCC ATATTTCCCT CAGCCTGCAT TTTGCCGATG CCCACCAGGG GTTATTGCAC	1680
	GGGAAGTCGG AGCTTGAGGC ACAATCTGTC GCGATCAGCC ATGGGCGACT GGTTGTGGCC	1740
	GATAGCGAAG GCAAGCTGTT TAGCGCCGCC ATTCCGAAGC AAGGGGATGG AAACGAACTG	1800
20	ARRATGRARG CCATGCCTCA GCATGCGCTC GATGRACATT TTGGTCATGA CCACCAGATT	1860
	TCTGGATTTT TCCATGACGA CCACGGCCAG CTTAATGCGC TGGTGAAAAA TAACTTCAGG	1920
25	CAGCAGCATG CCTGCCCGTT GGGTAACGAT CATCAGTTTC ACCCCGGCTG GAACCTGACT	1980
	GATGCOCTOG TTATCGACAA TCAGCTGGGG CTGCATCATA CCAATCCTGA ACCGCATGAG	2040
	ATTCTTGATA TGGGGCATTT AGGCAGCCTG GCGTTACAGG AGGGCAAGCT TCACTATTTT	2100
30	GACCAGCTGA CCAAAGGGTG GACTGGCGCG GAGTCAGATT GTAAGCAGCT GAAAAAAGGC	2160
	CTGGATGGAG CAGCTTATCT ACTGAAAGAC GGTGAAGTGA AACGCCTGAA TATTAATCAG	2220
35	AGCACCTCCT CTATCAAGCA CGGAACGGAA AACGTTTTTT CGCTGCCGCA TGTGCGCAAT	2280
	ARACCGGAGC CGGGAGATGC CCTGCAAGGG CTGAATAAMG ACGATAAGGC CCAGGCCATG	2340
	GCGGTGATTG GGGTAAATAA ATACCTGGCG CTGACGGAAA AAGGGGACAT TCGCTCCTTC	2400
40	CAGATAAAAC CCGGCACCCA GCAGTTGGAG CGGCCGGCAC AAACTCTCAG CCGCGAAGGT	2460
	ATCAGCGGG AACTGAAAGA CATTCATGTC GACCACAAGC AGAACCTGTA TGCCTTGACC	2520
45	CACGAGGGAG AGGTGTTTCA TCAGCCGCGT GAAGCCTGGC AGAATGGTGC CGAAAGCAGC	2580
	AGCTGGCACA AACTGGCGTT GCCACAGAGT GAAAGTAAGC TAAAAAGTCT GGACATGAGC	2640
	CATGAGCACA AACCGATTGC CACCITIGAA GACGGTAGCC AGCATCAGCT GAAGGCTGGC	2700
50	GGCTGGCACG CCTATGCGGC ACCTGAACGC GGGCCGCTGG CGGTGGGTAC CAGCGGTTCA	2760
	CARACCETCT TIRACCERCT RATECREGGE STERRAGGER AGGTGATCCC AGGCRGCGGG	2820
55	TTGACGGTTA AGCTCTCGGC TCAGACGGG GGAATGACCG GCGCCGAAGG GCGCAAGGTC	2880
"	AGCAGTAAAT TTTCCGAAAG GATCCGCGCC TATGCGTTCA ACCCAACAAT GTCCACGCCG	
	CGACCGATTA AAAATGCTGC TTATGCCACA CAGCACGGCT GGCAGGGGCG TGAGGGGTTG	
60	AAGCCGTTGT ACGAGATGCA GGGAGCGCTG ATTAAACAAC TGGATGCGCA TAACGTTCGT	
	CATARCECE CACAGCCAGA TITGCAGAGC ARACTGGARA CTCTGGATTT AGGCGRACAT	
C.E	CANADORD TOTAL CANADAGOOC TROOCCARCE AACTGGACCA GAGTGCAACC	

	. "	
	CGTTCGGTGA CCGTTTTAGG TCAACATCAG GGAGTGCTAA AAAGCAACGG TGAAATCAAT	3240
_	AGCGAATITA AGCCATCGCC CGGCAAGGCG TTGGTCCAGA GCTTTAACGT CAATCGCTCT	3300
5	GGTCAGGATC TAAGCAAGTC ACTGCAACAG GCAGTACATG CCACGCCGCC ATCCGCAGAG	3360
	AGTARACTGC AATCCATGCT GGGGCACTTT GTCAGTGCCG GGGTGGATAT GAGTCATCAG	3420
10	AAGGGCGAGA TCCCGCTGGG CCGCCAGCGC GATCCGAATG ATAAAACCGC ACTGACCAAA	3480
	TOGOGITTAA TITTAGATAC CGTGACCATC GGTGAACTGC ATGAACTGGC CGATAAGGCG	3540
	ARACTOGIAT CIGACCATAR ACCOGRIGCO GATCAGATAR ARCAGCIGCO CCAGCAGITC	3600
15	GATACGCTGC GTGARARGCG GTATGAGRGC RATCCGGTGR AGCRTTACAC CGATATGGGC	3660
•	TTCACCCATA ATAAGGCGCT GGAAGCAAAC TATGATGCGG TCAAAGCCTT TATCAATGCC	3720
20	TITAAGAAAG AGCACCACGG CGTCAATCTG ACCACGCGTA CCGTACTGGA ATCACAGGGC	3780
	AGTGCGGAGC TGGCGAAGAA GCTCAAGAAT ACGCTGTTGT CCCTGGACAG TGGTGAAAGT	3840
0.5	ATGAGCTTCA GCCGGTCATA TGGCGGGGGC GTCAGCACTG TCTTTGTGCC TACCCTTAGC	3900
25	AAGAAGGTGC CAGTTCCGGT GATCCCCGGA GCCGGCATCA CGCTGGATCG CGCCTATAAC	3960
	CTGAGCTTCA GTCGTACCAG CGGCGGATTG AACGTCAGTT TTGGCCGCGA CGGCGGGGTG	4020
30	AGTGGTAACA TCATGGTCGC TACCGGCCAT GATGTGATGC CCTATATGAC CGGTAAGAAA	4080
	ACCAGTGCAG GTAACGCCAG TGACTGGTTG AGCGCAAAAC ATAAAATCAG CCCGGACTTG	4140
	COTATCOGCO CTGCTGTGAG TGGCACCCTG CAAGGAACGC TACAAAACAG CCTGAAGTTT	4200
35	AAGCTGACAG AGGATGAGCT GCCTGGCTTT ATCCATGGCT TGACGCATGG CACGTTGACC	4260
	CCGGCAGAAC TGTTGCAAAA GGGGATCGAA CATCAGATGA AGCAGGGCAG CAAACTGACG	4320
40	TTTAGGGTCG ATACCTCGGC AAATCTGGAT CTGCGTGCCG GTATCAATCT GAACGAAGAC	4380
	GGCAGTAAAC CAAATGGTGT CACTGCCCGT GTTTCTGCCG GGCTAAGTGC ATCGGCAAAC	4440
	CTGGCCGCCG GCTCGCGTGA ACGCAGCACC ACCTCTGGCC AGTTTGGCAG CACGACTTCG	4500
45	GCCAGCAATA ACCGCCCAAC CTTCCTCAAC GGGGTCGGCG CGGGTGCTAA CCTGACGGCT	4560
	GCTTTAGGGG TTGCCCATTC ATCTACGCAT GAAGGGAAAC CGGTCGGGAT CTTCCCGGCA	4620
50	TITACCTOGA CCAATGITTC GGCAGCGCTG GCGCTGGATA ACCGTACCTC ACAGAGTATC	4680
	AGCCTGGAAT TGAAGCGCGC GGAGCCGGTG ACCAGCAACG ATATCAGCGA GTTGACCTCC	4740
	ACGCTGGGAA AACACTTTAA GGATAGCGCC ACAACGAAGA TGCTTGCCGC TCTCAAAGAG	4800
55	TTAGATGACG CTAAGCCCGC TGAACAACTG CATATTTTAC AGCAGCATTT CAGTGCAAAA	4860
	GATGTCGTCG GTGATGAACG CTACGAGGCG GTGCGCAACC TGAAAAAACT GGTGATACGT	4920
60	CAACAGGCTG CGGACAGCCA CAGCATGGAA TTAGGATCTG CCAGTCACAG CACGACCTAC	4980
	AATAATCTGT CGAGAATAAA TAATGACGGC ATTGTCGAGC TGCTACACAA ACATTTCGAT	5040
	GCGGCATTAC CAGCAAGCAG TGCCAAACGT CTTGGTGAAA TGATGAATAA CGATCCGGCA	5100
65		

	CTGAAAGATA	TTATTAAGCA	GCTGCAAAGT	ACCCCCTTCA	GCAGCGCCAG	CGTGTCGATG	5160
	GAGCTGAAAG	ATGGTCTGCG	TGAGCAGACG	GAAAAAGCAA	TACTGGACGG	TAAGGTCGGT	5220
5	CGTGAAGAAG	TGGGAGTACT	TTTCCAGGAT	CCTAACAACT	TGCGTGTTAA	ATCGGTCAGC	5280
	GTCAGTCAGT	CCGTCAGCAA	AAGCGAAGGC	TTCAATACCC	CAGCGCTGTT	ACTGGGGACG	5340
^	AGCAACAGCG	CTGCTATGAG	CATGGAGCGC	AACATCGGAA	CCATTAATTT	TAAATACGGC	5400
U	CAGGATCAGA	ACACOCCACG	GCGATTTACC	CTGGAGGGTG	GAATAGCTCA	GGCTAATCCG	5460
	CAGGTCGCAT	CTGCGCTTAC	TGATTTGAAG	DDDDAADDAA	TGGAAATGAA	GAGCTAA	551

This DNA molecule is known as the dspE gene for *Erwinia amylovora*. This isolated DNA molecule of the present invention encodes a protein or polypeptide which elicits a plant pathogen's hypersensitive response having an amino acid sequence of SEQ. ID. No. 8 as follows:

20 Met Glu Leu Lys Ser Leu Gly Thr Glu His Lys Ala Ala Val His Thr 1 5 10 15 Ala Ala His Asn Pro Val Gly His Gly Val Ala Leu Gln Gln Gly Ser 20 25 30 25 Ser Ser Ser Pro Gln Asn Ala Ala Ser Leu Ala Ala Glu Gly 35 40 45 Lys Asn Arg Gly Lys Met Pro Arg Ile His Gln Pro Ser Thr Ala Als 50 55 60 30 Asp Gly Ile Ser Ala Ala His Gln Gln Lys Lys Ser Phe Ser Leu Arg 65 70 75 80 35 Gly Cys Leu Gly Thr Lys Lys Fhe Ser Arg Ser Ala Pro Gln Gly Gln 85 90 95 Pro Gly Thr Thr His Ser Lys Gly Ala Thr Leu Arg Asp Leu Leu Ala 100 105 110 40 Arg Asp Asp Gly Glu Thr Gln His Glu Ala Ala Ala Pro Asp Ala Ala 115 120 125 Arg Leu Thr Arg Ser Gly Gly Val Lys Arg Arg Asn Met Asp Asp Met 130 135 140 45 Ala Gly Arg Pro Met Val Lys Gly Gly Ser Gly Glu Asp Lys Val Pro 145 150 160 50 Thr Gln Gln Lys Arg His Gln Leu Asn Asn Phe Gly Gln Met Arg Gln 165 170 175 Thr Met Leu Ser Lys Met Ala His Pro Ala Ser Ala Asn Ala Gly Asp 180 185 190 55 Arg Leu Gln His Ser Pro Pro His Ile Pro Gly Ser His His Glu Ile

		Glu '	Glu :	Pro '	Val (Gly	8er 7	Thr i	Ser	Lys	Ala :	hr 1	Thr A	la H	is Ai	la !	
5			Val	Glu :	Ile			31u .	Авр	Asp	Авр 235	Ber (3lu F	he G	ln G 2	ln 40	
		His	Gln		Arg 245	Leu	Ala i	Arg	Glu	Arg 250	Glu	Asn .	Pro I	ro G	ln P 55	LO	
10	Pro	Lys	Leu	Gly 260	Val	Ala	Thr	Pro	Ile 265	8er	Ala	Arg	Phe (31n F 270	ro I	iye	
15	Leu	Thr	Ala 275	Val	Ala	Glu	Ser	Val 280	Leu	Glu	Gly	Thr	Asp 1 285	Thr 7	fhr (3ln	
15		290					295				Gly	300					
20	305					310					1.ye 315					324	•
					325					330					335		
25				340					345	•	Ser			33V			
30			355					360			Phe		365				
		370					375				r His	380	ļ				
35	385					390)				395 r Gl:	,				40	
40					405	5				41	0 g Sex				413	•	
40				420)				42	5	o Gl		y Glu	esu Ser			
45			435	5			e His	440 Il:	0		a Pr	• GI	44: Let	•			•
	Glı	450 1 8e1)			s As	45! p Thi	5			ln Le	u Se	U			a A	
50	465 Gl ₃		s Let	и Туг				a As	p As	n A	47 rg Th 90		u Gl	n Ası	1 Le 49	u S	
55	Asj	ABA Ç	a Ly				u Ly	в Ье	u Va 50	al A	sp Ly	ns Il	e Ly	s 8e: 51:	r Ty		er
	۷a	l As _i				y G1	n Va	1 Al 52	a II		eu Tì	ır As	ap Th 52	r Pr		ly 3	ra
60	Hi	s Ly			r Il	e Me	t Pr 53	o Se		eu A	sp Al	a Se	•		u 86	er E	lis
65	11: 54:	e Se		u Se	r Le	u Hi 55	a Ph		a A	вр А	la H: 5:	ls G: 55	ln Gl	y Le	u Le	eu I	His 560

																_	
-	Gly				565					570	,				57.	•	
5	Leu	Val	val	Ala 580	qeA	Ser	Glu	Gly	Lys 585	Lev	. Phe	8ei	: Al	59	a Il O	e Pr	0
	Lys	Gln	Gly 595		GJĄ	Asn	Glu	Leu 600	Lye	Met	Ly	a Ale	60	t Pr 5	ro Gl	n Hi	.6
10	Ala	Leu 610		Glu	His	Phe	Gly 615	His	Ası	H1	e Gl	n Il 62	e Se O	r GI	y Pi	ne Ph	ıe
15	His 625		Asp	His	Gly	630	Leu	Ası	Al:	a Le	u Va 63	1 Ly 5	s As	n A	en P	be Ar	4 0
	Gln	Gli	His	ala	Cys 649	Pro	Lev	Gly	/ As:	n As 65	p Hi O	s Gl	n Pì	ie B	is P	ro G 55	ly
20	Trp) Ası	ı Let	Th:	ar (Ala	Lev	ı Va	1 Il 66	e As 5	ıp As	m Gl	n b	eu G 6	ly L 70	eu H	1 s
	Ris	Th	C Asi 67!		Gl:	ı Pro	Hi	G1: 68	u Il O	e Le	eu Ar	ap,∢ge	et G 6	ly B 85	is I	eu G	lly
25	Sez	Le: 69		a Let	ı Glı	n Gl	69:	γ Ι .γ 5	s Le	nı Hi	is T	YT P	ne A 00	ap G	iln I	beu 1	thr
·. 30	Lys 705		у Тт	p Th	r Gl	y Al: 71	a G1	u Se	r A	sp C	ys L 7	ys G 15	ln I	eu 1	Sya 1	iya (31y 720
	Lev	l As	p Gl	y Al	a Al 72	a Ty 5	r be	u Le	u L	78 A	вр G 30	ly G	lu V	al :	Був	Arg : 735	Leu
35	Ası	n Il	e As	n Gl 74	n Se O	r Th	r Se	r Se	er I:	le L 45	ys H	da G	lly ?	fhr (Glu 750	Asn	Val
	Ph	e Se	r Le 75		o Hi	.s Va	l Ar	g A	en L 50	ув Р	ro 0	3lu I	ore	31y 765	Asp	Ala	Leu
40	G1:		y L∈	eu Ae	m Ly	re As	77	np Li 15	ya A	la 6	In I	Ala !	Met . 780	Ala	Val	Ile	Gly
45	Va. 78		n Ly	/s T	r be	eu Al 79	La Le PO	ou T	hr G	ilu I	Lys (Gly 2 795	Asp	Ile	Arg	ser	Phe 800
	G1	n I	le L	ув Р	ro G: 80	ly Ti 05	nr G	Ln G	ln I	eu (Glu . 810	Arg	Pro	Ala	Gln	Thr 815	Leu
50	Se	r A	rg G	lu G 8:	ly I: 20	le S	er G	ly G	ilu 1	Geu : 825	Lys	Aep	Ile	His	Val 830	Asp	His
	Ľy	ns G		en L 35	eu T	YI A	la L	eu 1	hr 1	His	Glu	GΙΆ	Glu	Val 845	Phe	His	Gln
55	Pı		rg G 50	lu A	la T	rp G	ln A 8	sn (55	Ely :	Ala	Glu	Ser	8er 860	Ser	Tr	His	Lys
60		eu A	la L	eu P	ro G	ln s	er G 70	lu s	Ser	Lys	Leu	Lув 875	Ser	Leu	ABI	Met	880 880
-	H	is G	lu H	is L	ys P	ro I	le A	la:	Thr	Phe	890 890	Asp	Gly	8er	: Gli	Hi:	9 GL 5

	Leu Lys Ala Gly Gly Trp His Ala Tyr Ala Ala Pro Glu Arg Gly Pro 900 905 910
5	Leu Ala Val Gly Thr Ser Gly Ser Gln Thr Val Phe Asn Arg Leu Met 915 920 925
	Gln Gly Val Lys Gly Lys Val Ile Pro Gly Ser Gly Leu Thr Val Lys 930 935 940
10	Leu Ser Ala Gln Thr Gly Gly Met Thr Gly Ala Glu Gly Arg Lys Val 945 950 955 960
4	Ser Ser Lys Phe Ser Glu Arg Ile Arg Ala Tyr Ala Phe Asn Pro Thr 965 970 975
15	Met Ser Thr Pro Arg Pro Ile Lys Asn Ala Ala Tyr Ala Thr Gln His 980 985 990
20	Gly Trp Gln Gly Arg Glu Gly Leu Lys Pro Leu Tyr Glu Met Gln Gly 995 1000 1005
	Ala Leu Ile Lys Gln Leu Asp Ala His Asn Val Arg His Asn Ala Pro 1010 1015 1020
25	Gln Pro Asp Leu Gln Ser Lys Leu Glu Thr Leu Asp Leu Gly Glu His 1025 1030 1035 1040
	Gly Ala Glu Leu Leu Asn Asp Met Lys Arg Phe Arg Asp Glu Leu Glu 1045 1050 1055
30	Gln Ser Ala Thr Arg Ser Val Thr Val Leu Gly Gln His Gln Gly Val 1060 1065 1070
35	Leu Lys Ser Asn Gly Glu Ile Asn Ser Glu Phe Lys Pro Ser Pro Gly 1075 1080 1085
	Lys Ala Leu Val Gln Ser Phe Asn Val Asn Arg Ser Gly Gln Asp Leu 1090 1095 1100
40 .	Ser Lys Ser Leu Gln Gln Ala Val His Ala Thr Pro Pro Ser Ala Glu 1105 1110 1115 1120
45	Ser Lys Leu Gln Ser Met Leu Gly His Phe Val Ser Ala Gly Val Asp 1125 1130 1135
43	Net Ser His Gln Lys Gly Glu Ile Pro Leu Gly Arg Gln Arg Asp Pro 1140 1145 1150
50	Asn Asp Lys Thr Ala Leu Thr Lys Ser Arg Leu Ile Leu Asp Thr Val 1155 1160 1165
	Thr Ile Gly Glu Leu His Glu Leu Ala Asp Lys Ala Lys Leu Val Ser 1170 1175 1180
55	Asp His Lys Pro Asp Ala Asp Gln Ile Lys Gln Leu Arg Gln Gln Phe 1185 1190 1195 120
	Asp Thr Leu Arg Glu Lys Arg Tyr Glu Ser Asn Pro Val Lys His Tyr 1205 1210 1215
60	Thr Asp Met Gly Phe Thr His Asn Lys Ala Leu Glu Ala Asn Tyr Asp 1220 1225 1230
65	Ala Val Lys Ala Phe Ile Asn Ala Phe Lys Lys Glu His His Gly Val 1235 1240 1245

	Asn Leu Thr Thr Arg Thr Val Leu Glu Ser Gln Gly Ser Ala Glu Leu 1250 1255 1260
5	Ala Lys Lys Leu Lys Asn Thr Leu Leu Ser Leu Asp Ser Gly Glu Ser 1265 1270 1275 1280
4.0	Met Ser Phe Ser Arg Ser Tyr Gly Gly Val Ser Thr Val Phe Val 1285 1290 1295
10	Pro Thr Leu Ser Lys Lys Val Pro Val Pro Val Ile Pro Gly Ala Gly 1300 1305 1310
15	Ile Thr Leu Asp Arg Ala Tyr Asn Leu Ser Phe Ser Arg Thr Ser Gly 1315 1320 . 1325
	Gly Leu Asn Val Ser Phe Gly Arg Asp Gly Gly Val Ser Gly Asn Ile 1330 1335 1340
20	Met Val Ala Thr Gly His Asp Val Met Pro Tyr Met Thr Gly Lys 1345 1350 1355 1360
25	Thr Ser Ala Gly Asn Ala Ser Asp Trp Leu Ser Ala Lys His Lys Ile 1365 1370 1375
25	Ser Pro Amp Leu Arg Ile Gly Ala Ala Val Ser Gly Thr Leu Gln Gly 1380 1385 1390
30	Thr Leu Gln Asn Ser Leu Lys Phe Lys Leu Thr Glu Asp Glu Leu Pro 1395 1400 1405
	Gly Phe Ile His Gly Leu Thr His Gly Thr Leu Thr Pro Ala Glu Leu 1410 1415 1420
35	Leu Gln Lys Gly Ile Glu His Gln Met Lys Gln Gly Ser Lys Leu Thr 1425 1430 1435 1440
40	Phe Ser Val Asp Thr Ser Ala Asn Leu Asp Leu Arg Ala Gly Ile Asn 1445 1450 1455
40	Leu Asn Glu Asp Gly Ser Lys Pro Asn Gly Val Thr Ala Arg Val Ser 1460 1465 1470
45	Ala Gly Leu Ser Ala Ser Ala Asu Leu Ala Ala Gly Ser Arg Glu Arg 1475 1480 1485
•	Ser Thr Thr Ser Gly Gln Phe Gly Ser Thr Thr Ser Ala Ser Asn Asn 1490 1495 1500
50	Arg Pro Thr Phe Leu Asn Gly Val Gly Ala Gly Ala Asn Leu Thr Ala 1505 1510 1515 1520
	Ala Leu Gly Val Ala His Ser Ser Thr His Glu Gly Lys Pro Val Gly 1525 1530 1535
55	Ile Phe Pro Ala Phe Thr Ser Thr Asn Val Ser Ala Ala Leu Ala Leu 1540 1545 1550
60	Asp Asn Arg Thr Ser Gln Ser Ile Ser Leu Glu Leu Lys Arg Ala Glu 1555 1560 1565
	Pro Val Thr Ser Asn Asp Ile Ser Glu Leu Thr Ser Thr Leu Gly Lys 1570 1575 1580

	His Phe Lys Asp Ser Ala Thr Thr Lys Met Leu Ala Ala Leu Lys Glu 1585 1590 1595 1600
5	Leu Asp Asp Ala Lys Pro Ala Glu Gln Leu His Ile Leu Gln Gln His 1605 1610 1615
	Phe Ser Ala Lys Asp Val Val Gly Asp Glu Arg Tyr Glu Ala Val Arg 1620 1625 1630
10	Asn Leu Lys Lys Leu Val Ile Arg Gln Gln Ala Ala Asp Ser His Ser 1635 1640 1645
15	Met Glu Leu Gly Ser Ala Ser His Ser Thr Thr Tyr Asn Asn Leu Ser 1650 1655 1660
	Arg Ile Asn Asn Asp Gly Ile Val Glu Leu Leu His Lys His Phe Asp 1665 1670 1675 1680
20	Ala Ala Leu Pro Ala Ser Ser Ala Lys Arg Leu Gly Glu Met Met Asn 1685 1690 1695
	Asn Asp Pro Ala Leu Lys Asp Ile Ile Lys Gln Leu Gln Ser Thr Pro 1700 1705 1710
25	Phe Ser Ser Ala Ser Val Ser Met Glu Leu Lys Asp Gly Leu Arg Glu 1715 1720 1725
30	Gln Thr Glu Lys Ala Ile Leu Asp Gly Lys Val Gly Arg Glu Glu Val 1730 1735 · 1740
50	Gly Val Leu Phe Gln Asp Arg Asn Asn Leu Arg Val Lys Ser Val Ser 1745 1750 1755 1760
35	Val Ser Gln Ser Val Ser Lys Ser Glu Gly Phe Asn Thr Pro Ala Leu 1765 1770 1775
	Leu Leu Gly Thr Ser Asn Ser Ala Ala Met Ser Met Glu Arg Asn Ile 1780 1785 1790
40	Gly Thr Ile Asn Phe Lys Tyr Gly Gln Asp Gln Asn Thr Pro Arg Arg 1795 1800 1805
45	Phe Thr Leu Glu Gly Gly Ile Ala Gln Ala Asn Pro Gln Val Ala Ser 1810 1815 1820
	Ala Leu Thr Asp Leu Lys Cys Glu Gly Leu Glu Met Lys Ser 1825 1830 1835
50	This protein or polypeptide is about 198 kDa and has a pI of 8.98.
	The present invention relates to an isolated DNA molecule having a
	nucleotide sequence of SEQ. ID. No. 9 as follows:
55	ATGACATCGT CACAGCAGCG GGTTGAAAGG TTTTTACAGT ATTTCTCCGC CGGGTGTAAA 60
	ACGCCCATAC ATCTGAAAGA CGGGGTGTGC GCCCTGTATA ACGAACAAGA TGAGGAGGCG 120
~	GCGGTGCTGG AAGTACCGCA ACACAGCGAC AGCCTGTTAC TACACTGCCG AATCATTGAG 180
. 60	GCTGACCCAC ARACTTCRAT ARCCCTGTAT TCGATGCTAT TACAGCTGAA TTTTGARATG 240

GCGGCCATGC GCGGCTGTTG GCTGGCGCTG	GATGAACTGC	ACAACGTGCG	TTTATGITTT		300
CAGCAGTOGC TEGAGCATOT GGATGAAGCA	AGTITIAGCG	ATATCGTTAG	CGGCTTCATC	١	360
GAACATGCGG CAGAAGTGCG TGAGTATATA	GCGCAATTAG	ACGAGAGTAG	CGCGGCATAA		426

This is known as the dspF gene. This isolated DNA molecule of the present invention encodes a hypersensitive response elicitor protein or polypeptide having an amino acid sequence of SEQ. ID. No. 10 as follows:

40 This protein or polypeptide is about 16 kDa and has a pI of 4.45.

The hypersensitive response elicitor polypeptide or protein derived from *Pseudomonas syringae* has an amino acid sequence corresponding to SEQ. ID. No. 11 as follows:

•	Lys 65	Ser	Met	Ala	Ala	Авр (70	Gly :	Lув .	ala	Gly	Gly 75	GJÀ	Ile	Gl	u A	7 qa	Tal 30
	Ile	Ala	Ala	Leu	Asp 85	Lys	Leu	Ile	His	Glu 90	ГÀв	Leu	Gly	As	р А 9	sn 1 5	Phe
5	Gly	Ala	Ser	Ala 100	qaA	Ser	Ala	Ser	Gly 105	Thr	Gly	Gln	Glr	11	p L O	eu :	Met
	Thr	Gln	Val 115	Leu	Asn	Gly	Leu	Ala 120	ГÀв	Ser	Met	Leu	As; 12!	As 5	p I	eu	Leu
10	Thr	Lуя 130	Gln	Asp	Gly	Gly	Thr 135	Ser	Phe	Ser	Glu	Asp 140	Asj	p Me	et I	Pro	Met
•	145					150	Phe				155	i					160
	_				165		Trp			170)				•	173	
15				180					185	;				1	.90		Ile
	_		195					200					20	15			Gly
20		210					215	i					U				Ser .
	225					230	•				23	5					240
					245	5				25	0					25:	
25	Arg	Gly	, Leu	Glr 260		val	l Let	ı Ala	a G1 26	y G1	[D Y.	y Le	u G	ly '	Thr 270	Pro	o Val
			275	5				28	0				2	85			a Gln
30	Asp	290		e GJ1	a Let	u Le	29	y Gl: 5	у Le	u Le	eu Le	eu Ly 30	00 VB G	ly	Leu	G1	u Ala
	Th: 305		ı Lyı	a Asj	p Ala	a Gl; 31	y Gl:	n Th	r GJ	y Ti	or A	sp V: 15	al C	iln	Sez	: Se	r Ala 320
	Ala	a Gli	n Ile	e Al	a Th 32		u Le	u Va	1 86	2 T	br L 30	eu L	eu (lln	GJ?	7 Th 33	r Arg 15
35	Ası	a Gla	n Ala	a Al 34		a					•						

This hypersensitive response elicitor polypeptide or protein has a molecular weight of 34-35 kDa. It is rich in glycine (about 13.5%) and lacks cysteine and tyrosine. Further information about the hypersensitive response elicitor derived from Pseudomonas syringae is found in He, S. Y., H. C. Huang, and A. Collmer, "Pseudomonas syringae pv. syringae Harpings: a Protein that is Secreted via the Hrp Pathway and Elicits the Hypersensitive Response in Plants," Cell 73:1255-1266 (1993), which is hereby incorporated by reference. The DNA molecule encoding the hypersensitive response elicitor from Pseudomonas syringae has a nucleotide sequence corresponding to SEQ. ID. No. 12 as follows:

10 ATGCAGAGTC TCAGTCTTAA CAGCAGCTCG CTGCAAACCC CGGCAATGGC CCTTGTCCTG 60 GTACGTCCTG AAGCCGAGAC GACTGGCAGT ACGTCGAGCA AGGCGCTTCA GGAAGTTGTC 120 GTGAAGCTGG CCGAGGAACT GATGCGCAAT GGTCAACTCG ACGACAGCTC GCCATTGGGA 180 ARACTOTTOG CCARGTCORT GGCCGCAGAT GGCRAGGCGG GCGGCGGTAT TGRGGRTGTC 240 ATCGCTGCGC TGGACAAGCT GATCCATGAA AAGCTCGGTG ACAACTTCGG CGCGTCTGCG 300 15 AAGTCGATGC TCGATGATCT TCTGACCAAG CAGGATGGCG GGACAAGCTT CTCCGAAGAC 420 GATATGCCGA TGCTGAACAA GATCGCGCAG TTCATGGATG ACAATCCCGC ACAGTTTCCC 480 AAGCCGGACT CGGGCTCCTG GGTGAACGAA CTCAAGGAAG ACAACTTCCT TGATGGCGAC 540 GARACEGOTE COTTCCOTTC GECACTOGAC ATCATTEGEC AGCARCTEGE TARTCAGCAG 600 20 AGTGACGCTG GCAGTCTGGC AGGGACGGGT GGAGGTCTGG GCACTCCGAG CAGTTTTTCC 660 AACAACTOGT COGTGATGGG TGATCCGCTG ATCGACGCCA ATACCGGTCC CGGTGACAGC 720 GGCAATACCC GTGGTGAAGC GGGGCAACTG ATCGGCGAGC TTATCGACCG TGGCCTGCAA 780 TOGGTATTGG CCGGTGGTGG ACTGGGCACA CCCGTAAACA CCCCGCAGAC CGGTACGTCG 840 GCGAATGGCG GACAGTCCGC TCAGGATCTT GATCAGTTGC TGGGCGGCTT GCTGCTCAAG 900 25 GECCTGGAGG CAACGCTCAA GGATGCCGGG CAAACAGGCA CCGACGTGCA GTCGAGCGCT 960 GCGCAAATCG CCACCTTGCT GGTCAGTACG CTGCTGCAAG GCACCCGCAA TCAGGCTGCA 1020 1026 GCCTGA

Another potentially suitable hypersensitive response elicitor from Pseudomonas syringae is disclosed in U.S. Patent Application Serial No. 09/120,817, which is hereby incorporated by reference. The protein has a nucleotide sequence of SEQ. ID. No. 13 as follows:

	TCCACTTCGC TGATTITGAA ATTGGCAGAT TCATAGAAAC GTTCAGGTGT GGAAATCAGG	. 60
5	CTGAGTGCGC AGATTTCGTT GATAAGGGTG TGGTACTGGT CATTGTTGGT CATTTCAAGG	120
	CCTCTGAGTG CGGTGCGGAG CAATACCAGT CTTCCTGCTG GCGTGTGCAC ACTGAGTCGC	180
0	AGGCATAGGC ATTTCAGTTC CTTGCGTTGG TTGGGCATAT AAAAAAAGGA ACTTTTAAAA	240
	ACAGTGCAAT GAGATGCCGG CAAAACGGGA ACCGGTCGCT GCGCTTTGCC ACTCACTTCG	300
_	AGCAAGCTCA ACCCCAAACA TCCACATCCC TATCGAACGG ACAGCGATAC GGCCACTTGC	360
15	TCTGGTAAAC CCTGGAGCTG GCGTCGGTCC AATTGCCCAC TTAGCGAGGT AACGCAGCAT	420
	GAGCATCGGC ATCACACCCC GGCCGCAACA GACCACCACG CCACTCGATT TTTCGGCGCT	480
20	ARGCGGCARG AGTCCTCARC CARACACGTT CGGCGAGCAG ARCACTCAGC ARGCGATOGA	540
	CCCGAGTGCA CTGTTGTTCG GCAGCGACAC ACAGAAAGAC GTCAACTTCG GCACGCCCGA	600
~~	CAGCACCOTC CAGAATCCGC AGGACGCCAG CAAGCCCAAC GACAGCCAGT CCAACATCGC	660
25	TARATTORIC AGGCATTOR TCATGTCGTT GCTGCAGATG CTCACCAACT CCAATARAA	720
	GCAGGACACC AATCAGGAAC AGCCTGATAG CCAGGCTCCT TTCCAGAACA ACGGCGGGCT	780
30	COGTACACOG TCGGCCGATA GCGGGGGCGG CGGTACACCG GATGCGACAG GTGGCGGCGG	840
	CGGTGATACG CCAAGCGCAA CAGGCGGTGG CGGCGGTGAT ACTCCGACCG CAACAGGCGG	900
35	TEGOOGCAGC GETGGCGGCG GCACACCCAC TECAACAGGT GGCGGCAGCG GTGGCACACC	960
33	CACTGCAACA GGCGGTGGCG AGGGTGGCGT AACACCGCAA ATCACTCCGC AGTTGGCCAA	1020
	CCCTAACCGT ACCTCAGGTA CTGGCTCGGT GTCGGACACC GCAGGTTCTA CCGAGCAAGC	1080
40	COGCAAGATC AATGTGGTGA AAGACACCAT CAAGGTCGGC GCTGGCGAAG TCTTTGACGG	1140
	CCACGGCGCA ACCITCACIG CCGACAAATC TATGGGTAAC GGAGACCAGG GCGAAAATCA	1200
45	GAAGCCCATG TTCGAGCTGG CTGAAGGCGC TACGTTGAAG AATGTGAACC TGGGTGAGAA	1260
73	CGAGGTCGAT GGCATCCACG TGAAAGCCAA AAACGCTCAG GAAGTCACCA TTGACAACGT	1320
	GCATGCCCAG AACGTCGGTG AAGACCTGAT TACGGTCAAA GGCGAGGGAG GCGCAGCGGT	1380
50	CACTAATCTG AACATCAAGA ACAGCAGTGC CAAAGGTGCA GACGACAAGG ITGTCCAGCT	1440
	CAACGCCAAC ACTCACTTGA AAATCGACAA CTTCAAGGCC GACGATTTCG GCACGATGGT	1500
55	TOGCACCAAC GOTGGCAAGC AGITTGATGA CATGAGCATC GAGCTGAACG GCATCGAAGC	1560
,,,	TAACCACGGC AAGTTCGCCC TGGTGAAAAG CGACAGTGAC GATCTGAAGC TGGCAACGGG	1620
	CAACATCGCC ATGACCGACG TCAAACACGC CTACGATAAA ACCCAGGCAT CGACCCAACA	1680
60	CACCGAGGIT TGAATCCAGA CAAGTAGCIT GAAAAAAGGG GGTGGACIC	172

This DNA molecule is known as the dspE gene for *Pseudomonas syringae*. This isolated DNA molecule of the present invention encodes a protein or polypeptide which elicits a plant pathogen's hypersensitive response having an amino acid sequence of SEQ. ID. No. 14 as follows:

										•						
5	1				Ile 5					10					13	
10	Asp	Phe	Ser	Ala 20	Leu	Ser	Gly	Гув	Ser 25	Pro	Gln	Pro	Asn	Thr 30	Phe	Gly
	Glu	Gln	Asn 35	Thr	Gln	Gln	Ala	Ile 40	Asp	Pro	8er	Ala	Leu 45	Leu	Phe	GJÀ
15	Ser	Asp 50	Thr	Gln	Lys	qaA	Val 55	Asn	Phe	Gly	Thr	Pro 60	Asp	Ser	Thr	Val
_1	Gln 65	Asn	Pro	Gln	Asp	Ala 70	Ser	ŗå	Pro	Asn	Авр 75	5er	Gln	Ser	Ави	Ile 80
20	Ala	Гув	Leu	Ile	Ser 85	Ala	Leu	Ile	Met	Ser 90	Leu	Leu	Gln	Met	Leu 95	Thr
25	Asn	Ser	Asn	L ув 100		Gln	Asp	Thr	Asn 105	Gln	Glu	Gln	Pro	Asp qua	Ser	Gln
	Ala	Pro	Phe		Asn.	Asn	Gly	Gly 120	Leu	G1y	Thr	Pro	ser 125	Ala	Asp	Ser
30	Gly	Gly		Gly	Thr	Pro	Авр 135	Ala	Thr	Gly	Gly	Gly 140	Gly	Gly	, yaż	Thr
	Pro		Ala	Thr	Gly	Gly 150	Gly	G17	, Gl3	gaA v	155	Pro	Tha	Ale	a Thi	160
35	Gly	Gl	/ Gly	/ Sei	: Gly 165		Gl3	gl ₃	Th	170	Thi	. Ala	Thi	Gl ₃	7 Gl	e Gly
40	Sez	Gl _y	, Gl	Thi 180		Thi	- Ala	Th:	r Gl;	y Gl3 5	y Gly	y Gla	a Gl	y Gl; 19	y Va. O	l Thr
	Pro	Gl:	110 19		r Pro	Gli	Le:	Al: 20	а Ав: 0	n Pro	aA c	a Ar	g Th	r Se 5	r Gl	y Thr
45	Gly	/ Se:		l Se	r Asj	Th:	r Ala 21	a Gl	y Se	r Th	r Gl	u Gl 22	n Al O	a Gl	у гу	s Ile
	As: 22!		l Va	l Ly	в Авј	23	r Il	e Ly	s Va	1 G1	y Al 23	a Gl 5	y Gl	u Va	l Ph	e Asp 240
50	Gl	tH v	s Gl	y Al	a Thi		e Th	r Al	a As	р Ly 25	ទ Se 0	r Me	t Gl	у Ав	n Gl 25	y Asp is
55	Gli	n G1	y Gl	u As: 26		r FÀ	s Pr	o Me	t Pb 26	e Gl	n Pe	u Al	a Gl	u G1 27	LA Y.	a Thr

	Leu	Lys	Asn 275	Val	Asn	Leu	Gly	Glu 280	Asn	Glu	val	qaA	Gly 285	Ile	His	Val
5	Lув	Ala 290		Asn	Ala	Gln	Glu 295	Val	Thr	Ile	Ąsp	200 200	Val	His	Ala	Gln
_	Asn 305	Val	Gly	Glu	qaA	Leu 310	Ile	Thr	Val	Lys	Gly 315	Glu	Gly	Gly	Ala	Ala 320
10	Val	Thr	Asn	Leu	Asn 325		Lys	Asn	Ser	8er 330	Ala	ГÀВ	Gly	Ala	Авр 335	Asp
15	Гув	Val	Val	Gln 340		Asn	Ala	Asn	Thr 345	His	Leu	Lys	Ile	Авр 350	Asn	Phe
	Lys	Ala	Авр 355		Phe	Gly	Thr	Met 360	Val	Arg	Thr	Asr	Gly 365	Gly	Гув	Gln
20	Phe	Asp 370		Met	Ser	Ile	375	Leu	Asn	a Gly	Ile	380	ı Ala	Авт	His	Gly
	Lys 385		. Ala	Leu	Val	. Lys 390	Ser	Asp	Ser	. Yel	395	Let 5	ı Lyı	. Lev	ı Ala	Thr 400
25 ⁻	Gly	Asi	ıle	: Ala	Met 405		r Asy	val	. Ly	8 His 410	s Ala	а Ту	r Asj	p Lyı	41!	r Gln
30	Ala	s Sez	Thr	Gl: 420		Th:	r Glı	ı Let	ı							

This protein or polypeptide is about 42.9 kDa.

This hypersensitive response elicitor from *Pseudomonas syringae* has 1

35 hypersensitive response eliciting domain. This domain extends, within SEQ. ID. No. 14, from amino acid 45 to amino acid 102, particularly from amino acid 58 to amino acid 92. The acidic unit in the first domain extends, within SEQ. ID. No. 14, from amino acid 45 to amino acid 79, particularly from amino acid 58 to amino acid 79. The alpha-helix in the first domain extends, within SEQ. ID. No. 14, from amino acid 79 to amino acid 102, particularly from amino acid 79 to amino acid 92.

The hypersensitive response elicitor polypeptide or protein derived from *Pseudomonas solanacearum* has an amino acid sequence corresponding to SEQ. ID. No. 15 as follows:

Met Ser Val Gly Asn Ile Gln Ser Pro Ser Asn Leu Pro Gly Leu Gln
1 5 10 15

Asn Leu Asn Leu Asn Thr Asn Thr Asn Ser Gln Gln Ser Gly Gln Ser

	Val	Gln	Двр 35	Leu	Ile	Гув	Gln	Val 40	Glu	Lys	Авр	Ile	Leu 45	L As:	n I	le I	(le	
	Ala	Ala 50	•	Val	Gln	Lys	Ala 55	Ala	Gln	Ser	Ala	Gly 60	Gl	7 As	n T	hr (31y	•
5	Asn 65		Gly	Asn	Ala	Pro 70		Гув	Дар	Gly	Asn 75	Ala	Asi	n Al	a G	ily :	Ala 80	L
		Авр	Pro	Ser	Ьув 85		Авр	Pro	Ser	Був 90	Ser	Gli	Al.	a Pı	eo 6	31n 95	Sei	r
10	Ala	Asn	Lys	Thr 100	Gly	Asn	Val	Asp	Asp 105	Ala	. Asr	Ası	ı Gl	n A	sp 1	Pro	Mei	t
	Gln	Ala	Leu 115		Gln	Leu	Leu	Glu 120	Asp	Let	ı Val	Ly	s Le	u L 15	eu :	Lys	Al	a
	Ala	Leu 130		Met	Gln	Glr	135		Gl)	r Ası	a Asj	14	s G) 0	ly A	sn :	GJY	۷a	1
15	Gly 145		Ala	Asn	Gly	Ala 150	Lys)	Gly	, Ali	a Gl	y Gl; 15	5 5	n G	ly G	ly	Leu	Al 16	.a. i 0
	Glu	Ala	Lev	Gln	165		e Glı	ı Gli	a Ile	е Le [.] 17	u Al: O	a Gl	n L	eu C	ly	Gly 175	G]	Y
20				180)				18	5	y Va			•	190			
			19!	5				20	0		a Gl		4	.05				
		210)				21	5			n Al	2:	20					
25	225	5				23	0				la As 23	15					4	.40
					24	5				2	78 I. 50					~3	3	
30				26	0				20	55	ly L				2/(,		
			27	5				2	BO		sn A			285				
		29	0				2:	95			er A	-	300					
35	30	5				3	10					15						341
	Va	1 Gl	n Il	e Le	nu G] 32	ln G 15	ln M	et L	eu A	la A	la G	ln i	Asn	Gly	GJ.	y S:	er 35	Gli

Gln Ser Thr Ser Thr Gln Pro Met

It is encoded by a DNA molecule having a nucleotide sequence corresponding SEQ. ID. No. 16 as follows:

_	ATGTCAGTCG GARACATCCA GAGCCCGTCG AACCTCCCGG GTCTGCAGAA CCTGAACCTC	60
5		-
	AACACCAACA CCAACAGCCA GCAATOGGGC CAGTCOGTGC AAGACCTGAT CAAGCAGGTC	120
	GAGAAGGACA TCCTCAACAT CATCGCAGCC CTCGTGCAGA AGGCCGCACA GTCGGCGGGC	180
	GGCAACACCG GTAACACCGG CAACGCGCCG GCGAAGGACG GCAATGCCAA CGCGGGCGCC	240
•	AACGACCOGA GCAAGAACGA CCCGAGCAAG AGCCAGGCTC CGCAGTCGGC CAACAAGACC	300
. 10	GGCAACGTCG ACGACGCCAA CAACCAGGAT CCGATGCAAG CGCTGATGCA GCTGCTGGAA	360
	GACCTGGTGA AGCTGCTGAA GGCGGCCCTG CACATGCAGC AGCCCGGCGG CAATGACAAG	420
	GGCAACGGCG TGGGCGGTGC CAACGGCGCC AAGGGTGCCG GCGGCCAGGG CGGCCTGGCC	480
	GAAGOSCTGC AGGAGATOGA GCAGATOCTC GCCCAGCTCG GCGGCGGCGG TGCTGGCGCC	540
	GECGCCCCG GTGGCGGTGT CGGCGGTGCT GGTGCCGCGG ATGGCGGCTC CGGTGCGGGT	600
15	GGCGCAGGCG GTGCGAACGG CGCCGACGGC GGCAATGGCG TGAACGGCAA CCAGGCGAAC	660
	GGCCCGCAGA ACGCAGGCGA TGTCAACGGT GCCAACGGCG CGGATGACGG CAGCGAAGAC	720
	CAGGGCGGCC TCACCGGCGT GCTGCAAAAG CTGATGAAGA TCCTGAACGC GCTGGTGCAG	780
	ATGATGCAGC AAGGCGGCCT CGGCGGCGGC AACCAGGCGC AGGGCGGCTC GAAGGGTGCC	840
	GGCAACGCCT CGCCGGCTTC CGGCGCGAAC CCGGGCGCGA ACCAGCCCGG TTCGGCGGAT	900
20	GATCAATCGT CCGGCCAGAA CAATCTGCAA TCCCAGATCA TGGATGTGGT GAAGGAGGTC	96
	GTCCAGATCC TGCAGCAGAT GCTGGCGGCG CAGAACGGCG GCAGCCAGCA GTCCACCTCG	102
	ACGCAGCCGA TGTAA	103

25 Further information regarding the hypersensitive response elicitor polypeptide or protein derived from Pseudomonas solanacearum is set forth in Arlat, M., F. Van Gijsegem, J. C. Huet, J. C. Pemollet, and C. A. Boucher, "PopA1, a Protein which Induces a Hypersensitive-like Response in Specific Petunia Genotypes, is Secreted via the Hrp Pathway of Pseudomonas solanacearum," EMBO J. 13:543-533 (1994), which is hereby incorporated by reference.

The hypersensitive response elicitor from *Pseudomonas solanacearum* has 2 hypersensitive response eliciting domains. The first domain extends, within SEQ. ID.

No. 15, from amino acid 85 to amino acid 131, particularly from amino acid 95 to amino acid 123. The acidic unit in the first domain extends, within SEQ. ID. No. 15, from amino acid 85 to amino acid 111, particularly from amino acid 95 to amino acid 123. The alpha-helix in the first domain extends, within SEQ. ID. No. 15, from amino acid 85 to amino acid 111, particularly from amino acid 95 to amino acid 111. The second domain extends, within SEQ. ID. No. 15, from amino acid 195 to amino acid 264, particularly from amino acid 229 to amino acid 258. The acidic unit in the second domain extends, within SEQ. ID. No. 15, from amino acid 195 to amino acid 246, particularly from amino acid 229 to amino acid 264. The alpha-helix in the second domain extends, within SEQ. ID. No. 15, from amino acid 246 to amino acid 264, particularly from amino acid 246 to amino acid 258.

The N-terminus of the hypersensitive response elicitor polypeptide or protein from *Xanthomonas campestris* has an amino acid sequence corresponding to SEO. ID. No. 17 as follows:

- Met Asp Gly Ile Gly Asn His Phe Ser Asn
 1 5 10
- The hypersensitive response elicitor polypeptide or protein from Xanthomonas campestris pv. pelargonii is heat stable, protease sensitive, and has a molecular weight of 20 kDa. It includes an amino acid sequence corresponding to SEQ. ID. No. 18 as follows:
- 25 Ser Ser Gln Gln Ser Pro Ser Ala Gly Ser Glu Gln Gln Leu Asp Gln
 1 5 10 15

 Leu Leu Ala Met
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Isolation of Erwinia carotovora hypersensitive response elictor protein or polypeptide is described in Cui et al., "The RsmA Mutants of Erwinia carotovora subsp. carotovora Strain Ecc71 Overexpress hrp N_{Ecc} and Elicit a Hypersensitive Reaction-like Response in Tobacco Leaves," MPMI, 9(7):565-73 (1996), which is hereby incorporated by reference. The hypersensitive response elicitor protein or polypeptide of Erwinia stewartii is set forth in Ahmad et al., "Harpin is Not

Necessary for the Pathogenicity of Erwinia stewartii on Maize," 8th Int'l. Cong.

Molec. Plant-Microbe Interact., July 14-19, 1996 and Ahmad, et al., "Harpin is Not Necessary for the Pathogenicity of Erwinia stewartii on Maize," Am. Mtg. Am.

Phytopath. Soc., July 27-31, 1996, which are hereby incorporated by reference.

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Hypersensitive response elicitor proteins or polypeptides from
Phytophthora parasitica, Phytophthora cryptogea, Phytophthora cinnamoni,
Phytophthora capsici, Phytophthora megasperma, and Phytophthora citrophthora are
described in Kaman, et al., "Extracellular Protein Elicitors from Phytophthora: Most
Specificity and Induction of Resistance to Bacterial and Fungal Phytopathogens,"
Molec. Plant-Microbe Interact., 6(1):15-25 (1993), Ricci et al., "Structure and
Activity of Proteins from Pathogenic Fungi Phytophthora Eliciting Necrosis and
Accepted Resistance in Tobacco," Eur. I. Biochem., 183:555-63 (1989), Ricci et al.,

Activity of Proteins from Pathogenic Fungi Phytophthora Eliciting Necrosis and Acquired Resistance in Tobacco," <u>Eur. J. Biochem.</u>, 183:555-63 (1989), Ricci et al., "Differential Production of Parasiticein, and Elicitor of Necrosis and Resistance in Tobacco, by Isolates of Phytophthora parasitica," <u>Plant Path.</u> 41:298-307 (1992),

Baillreul et al, "A New Elicitor of the Hypersensitive Response in Tobacco: A Fungal Glycoprotein Elicits Cell Death, Expression of Defence Genes, Production of Salicylic Acid, and Induction of Systemic Acquired Resistance," Plant J., 8(4):551-60 (1995), and Bonnet et al., "Acquired Resistance Triggered by Elicitors in Tobacco and Other Plants," Eur. J. Plant Path., 102:181-92 (1996), which are hereby incorporated by reference.

Another hypersensitive response elicitor in accordance with the present invention is from *Clavibacter michiganensis* subsp. sepedonicus which is fully described in U.S. Patent Application Serial No. 09/136,625, which is hereby incorporated by reference.

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The above elicitors are exemplary. Other elicitors can be identified by growing fungi or bacteria that elicit a hypersensitive response under conditions which genes encoding an elicitor are expressed. Cell-free preparations from culture supernatants can be tested for elicitor activity (i.e. local necrosis) by using them to infiltrate appropriate plant tissues.

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Fragments of the above hypersensitive response elicitor polypeptides or proteins as well as fragments of full length elicitors from other pathogens are encompassed by the method of the present invention.

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Suitable fragments can be produced by several means. In the first, subclones of the gene encoding a known elicitor protein are produced by conventional molecular genetic manipulation by subcloning gene fragments. The subclones then are expressed in vitro or in vivo in bacterial cells to yield a smaller protein or peptide that can be tested for elicitor activity according to the procedure described below.

As an alternative, fragments of an elicitor protein can be produced by digestion of a full-length elicitor protein with proteolytic enzymes like chymotrypsin or *Staphylococcus* proteinase A, or trypsin. Different proteolytic enzymes are likely to cleave elicitor proteins at different sites based on the amino acid sequence of the elicitor protein. Some of the fragments that result from proteolysis may be active elicitors of resistance.

In another approach, based on knowledge of the primary structure of the protein, fragments of the elicitor protein gene may be synthesized by using the PCR technique together with specific sets of primers chosen to represent particular portions of the protein. These then would be cloned into an appropriate vector for expression of a truncated peptide or protein.

Chemical synthesis can also be used to make suitable fragments. Such a synthesis is carried out using known amino acid sequences for the elicitor being produced. Alternatively, subjecting a full length elicitor to high temperatures and pressures will produce fragments. These fragments can then be separated by conventional procedures (e.g., chromatography, SDS-PAGE).

An example of suitable fragments of a hypersensitive response elicitor which do elicit a hypersensitive response are *Erwinia amylovora* fragments including a C-terminal fragment of the amino acid sequence of SEQ. ID. No. 3, an N-terminal fragment of the amino acid sequence of SEQ. ID. No. 3, or an internal fragment of the amino acid sequence of SEQ. ID. No. 3. The C-terminal fragment of the amino acid sequence of SEQ. ID. No. 3 can span amino acids 105 and 403 of SEQ. ID. No. 3. The N-terminal fragment of the amino acid sequence of SEQ. ID. No. 3 can span the following amino acids of SEQ. ID. No. 3: 1 and 98, 1 and 104, 1 and 122, 1 and 168, 1 and 218, 1 and 266, 1 and 342, 1 and 321, and 1 and 372. The internal fragment of the amino acid sequence of SEQ. ID. No. 3 can span the following amino acids of

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SEQ. ID. No. 3: 76 and 209, 105 and 209, 99 and 209, 137 and 204, 137 and 200, 109 and 204, 109 and 200, 137 and 180, and 105 and 180.

Suitable DNA molecules are those that hybridize to the DNA molecule comprising a nucleotide sequence of SEQ. ID. Nos. 2, 4, 5, 7, 9, 12, 13, and 16 under stringent conditions. An example of suitable high stringency conditions is when hybridization is carried out at 65°C for 20 hours in a medium containing 1M NaCl, 50 mM Tris-HCl, pH 7.4, 10 mM EDTA, 0.1% sodium dodecyl sulfate, 0.2% ficoll, 0.2% polyvinylpyrrolidone, 0.2% bovine serum albumin, 50 μm g/ml E. coli DNA. Suitable stringency conditions also include hybridization in a hybridization buffer comprising 0.9M sodium citrate ("SSC") buffer at a temperature of 37°C where 10 . hybridized nucleic acids remain bound when subject to washing the SSC buffer at a temperature of 37°C; and preferably in a hybridization buffer comprising 20% formamide in 0.9M SSC buffer at a temperature of 42°C where hybridized nucleic acids remain bound when subject to washing at 42°C with 0.2x SSC buffer at 42°C.

Variants may be made by, for example, the deletion or addition of amino acids that have minimal influence on the properties, secondary structure and hydropathic nature of the polypeptide. For example, a polypeptide may be conjugated to a signal (or leader) sequence at the N-terminal end of the protein which cotranslationally or post-translationally directs transfer of the protein. The polypeptide may also be conjugated to a linker or other sequence for ease of synthesis, 20 : purification, or identification of the polypeptide.

A particularly advantageous aspect of the present invention involves utilizing a protein having a pair or more, particularly 3 or more, coupled domains. These domains can be from different source organisms. When a DNA molecule encoding such a protein is prepared, it can be advantageously used to make transgenic plants. The use of a gene encoding such domains, as opposed to a gene encoding a full length hypersensitive response elicitor, has a number of benefits. Firstly, such a gene is easier to synthesize. More significantly, the use of a plurality of domains together from different source organisms can impart their combined benefits to a transgenic plant.

The DNA molecule encoding the hypersensitive response elicitor polypeptide or protein can be incorporated in cells using conventional recombinant

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DNA technology. Generally, this involves inserting the DNA molecule into an expression system to which the DNA molecule is heterologous (i.e. not normally present). The heterologous DNA molecule is inserted into the expression system or vector in proper sense orientation and correct reading frame. The vector contains the necessary elements for the transcription and translation of the inserted protein-coding sequences.

U.S. Patent No. 4,237,224 to Cohen and Boyer, which is hereby incorporated by reference, describes the production of expression systems in the form of recombinant plasmids using restriction enzyme cleavage and ligation with DNA ligase. These recombinant plasmids are then introduced by means of transformation and replicated in unicellular cultures including procaryotic organisms and eucaryotic cells grown in tissue culture.

Recombinant genes may also be introduced into viruses, such as vaccina virus. Recombinant viruses can be generated by transfection of plasmids into cells infected with virus.

Suitable vectors include, but are not limited to, the following viral vectors such as lambda vector system gt11, gt WES.tB, Charon 4, and plasmid vectors such as pBR322, pBR325, pACYC177, pACYC1084, pUC8, pUC9, pUC18, pUC19, pLG339, pR290, pKC37, pKC101, SV 40, pBluescript II SK +/- or KS +/- (see "Stratagene Cloning Systems" Catalog (1993) from Stratagene, La Jolla, Calif, which is hereby incorporated by reference), pQE, pIH821, pGEX, pET series (see F.W. Studier et. al., "Use of T7 RNA Polymerase to Direct Expression of Cloned Genes," Gene Expression Technology vol. 185 (1990), which is hereby incorporated by reference), and any derivatives thereof. Recombinant molecules can be introduced into cells via transformation, particularly transduction, conjugation, mobilization, or electroporation. The DNA sequences are cloned into the vector using standard cloning procedures in the art, as described by Sambrook et al., Molecular Cloning: A Laboratory Manual. Cold Springs Laboratory, Cold Springs Harbor, New York (1989), which is hereby incorporated by reference.

A variety of host-vector systems may be utilized to express the proteinencoding sequence(s). Primarily, the vector system must be compatible with the host cell used. Host-vector systems include but are not limited to the following: bacteria

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transformed with bacteriophage DNA, plasmid DNA, or cosmid DNA; microorganisms such as yeast containing yeast vectors; mammalian cell systems infected with virus (e.g., vaccinia virus, adenovirus, etc.); insect cell systems infected with virus (e.g., baculovirus); and plant cells infected by bacteria. The expression elements of these vectors vary in their strength and specificities. Depending upon the host-vector system utilized, any one of a number of suitable transcription and translation elements can be used.

Different genetic signals and processing events control many levels of gene expression (e.g., DNA transcription and messenger RNA (mRNA) translation).

Transcription of DNA is dependent upon the presence of a promotor which is a DNA sequence that directs the binding of RNA polymerase and thereby promotes mRNA synthesis. The DNA sequences of eucaryotic promotors differ from those of procaryotic promotors. Furthermore, eucaryotic promotors and accompanying genetic signals may not be recognized in or may not function in a procaryotic system, and, further, procaryotic promotors are not recognized and do not function in eucaryotic cells.

Similarly, translation of mRNA in procaryotes depends upon the presence of the proper procaryotic signals which differ from those of eucaryotes. Efficient translation of mRNA in procaryotes requires a ribosome binding site called the Shine-Dalgarno ("SD") sequence on the mRNA. This sequence is a short nucleotide sequence of mRNA that is located before the start codon, usually AUG, which encodes the amino-terminal methionine of the protein. The SD sequences are complementary to the 3'-end of the 16S rRNA (ribosomal RNA) and probably promote binding of mRNA to ribosomes by duplexing with the rRNA to allow correct positioning of the ribosome. For a review on maximizing gene expression, see Roberts and Lauer, Methods in Enzymology, 68:473 (1979), which is hereby incorporated by reference.

Promotors vary in their "strength" (i.e. their ability to promote transcription). For the purposes of expressing a cloned gene, it is desirable to use strong promotors in order to obtain a high level of transcription and, hence, expression of the gene. Depending upon the host cell system utilized, any one of a number of suitable promotors may be used. For instance, when cloning in E. coli, its

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bacteriophages, or plasmids, promotors such as the T7 phage promoter, *lac* promotor, *trp* promotor, *rec*A promotor, ribosomal RNA promotor, the P_R and P_L promotors of coliphage lambda and others, including but not limited, to *lac*UV5, *omp*F, *bla*, *lpp*, and the like, may be used to direct high levels of transcription of adjacent DNA segments. Additionally, a hybrid *trp-lac*UV5 (*tac*) promotor or other *E. coli* promotors produced by recombinant DNA or other synthetic DNA techniques may be used to provide for transcription of the inserted gene.

Bacterial host cell strains and expression vectors may be chosen which inhibit the action of the promotor unless specifically induced. In certain operations, the addition of specific inducers is necessary for efficient transcription of the inserted DNA. For example, the *lac* operon is induced by the addition of lactose or IPTG (isopropylthio-beta-D-galactoside). A variety of other operons, such as *trp*, *pro*, etc., are under different controls.

transcription and translation in procaryotic cells. These transcription and translation initiation signals may vary in "strength" as measured by the quantity of gene specific messenger RNA and protein synthesized, respectively. The DNA expression vector, which contains a promotor, may also contain any combination of various "strong" transcription and/or translation initiation signals. For instance, efficient translation in E. coli requires an SD sequence about 7-9 bases 5' to the initiation codon ("ATG") to provide a ribosome binding site. Thus, any SD-ATG combination that can be utilized by host cell ribosomes may be employed. Such combinations include but are not limited to the SD-ATG combination from the cro gene or the N gene of coliphage lambda, or from the E. coli tryptophan E, D, C, B or A genes. Additionally, any SD-ATG combination produced by recombinant DNA or other techniques involving incorporation of synthetic nucleotides may be used.

Once the isolated DNA molecule encoding the hypersensitive response elicitor polypeptide or protein has been cloned into an expression system, it is ready to be incorporated into a host cell. Such incorporation can be carried out by the various forms of transformation noted above, depending upon the vector/host cell system. Suitable host cells include, but are not limited to, plant cells as well as

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prokaryotic and eukaryotic cells, such as bacteria, virus, yeast, mammalian, insect cells, and the like.

The present invention further relates to methods of imparting disease resistance to plants, enhancing plant growth, effecting insect control and/or imparting stress resistance to plants. These methods involve applying a hypersensitive response elicitor polypeptide or protein to all or part of a plant or a plant seed under conditions where the polypeptide or protein contacts all or part of the cells of the plant or plant seed. Alternatively, the hypersensitive response elicitor protein or polypeptide can be applied to plants such that seeds recovered from such plants themselves are able to impart disease resistance in plants, to enhance plant growth, to effect insect control, and/or to impart stress resistance.

As an alternative to applying a hypersensitive response elicitor polypeptide or protein to plants or plant seeds in order to impart disease resistance in plants, to effect plant growth, to control insects, and/or to impart stress resistance to the plants or plants grown from the seeds, transgenic plants or plant seeds can be utilized. When utilizing transgenic plants, this involves providing a transgenic plant transformed with a DNA molecule encoding a hypersensitive response elicitor polypeptide or protein and growing the plant under conditions effective to permit that DNA molecule to impart disease resistance to plants, to enhance plant growth, to control insects, and/or to impart stress resistance. Alternatively, a transgenic plant seed transformed with a DNA molecule encoding a hypersensitive response elicitor polypeptide or protein can be provided and planted in soil. A plant is then propagated from the planted seed under conditions effective to permit that DNA molecule to impart disease resistance to plants, to enhance plant growth, to control insects, and/or to impart stress resistance.

The method of the present invention can be utilized to treat a wide variety of plants or their seeds to impart disease resistance, enhance growth, control insects, and/or to impart stress resistance. Suitable plants include dicots and monocots. More particularly, useful crop plants can include: alfalfa, rice, wheat, barley, rye, cotton, sunflower, peanut, corn, potato, sweet potato, bean, pea, chicory, lettuce, endive, cabbage, brussel sprout, beet, parsnip, turnip, cauliflower, broccoli, turnip, radish, spinach, onion, garlic, eggplant, pepper, celery, carrot, squash,

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pumpkin, zucchini, cucumber, apple, pear, melon, citrus, strawberry, grape, raspberry, pineapple, soybean, tobacco, tomato, sorghum, and sugarcane. Examples of suitable ornamental plants are: Arabidopsis thaliana, Saintpaulia, petunia, pelargonium, poinsettia, chrysanthemum, carnation, and zinnia.

With regard to the use of the hypersensitive response elicitor protein or polypeptide of the present invention in imparting disease resistance, absolute immunity against infection may not be conferred, but the severity of the disease is reduced and symptom development is delayed. Lesion number, lesion size, and extent of sporulation of fungal pathogens are all decreased. This method of imparting disease resistance has the potential for treating previously untreatable diseases, treating diseases systemically which might not be treated separately due to cost, and avoiding the use of infectious agents or environmentally harmful materials.

The method of imparting pathogen resistance to plants in accordance with the present invention is useful in imparting resistance to a wide variety of pathogens including viruses, bacteria, and fungi. Resistance, inter alia, to the following viruses can be achieved by the method of the present invention: Tobacco mosaic virus and Tomato mosaic virus. Resistance, inter alia, to the following bacteria can also be imparted to plants in accordance with present invention:

Pseudomonas solancearum, Pseudomonas syringae pv. tabaci, and Xanthamonas campestris pv. pelargonii. Plants can be made resistant, inter alia, to the following fungi by use of the method of the present invention: Fusarium oxysporum and Phytophthora infestans.

With regard to the use of the hypersensitive response elicitor protein or polypeptide of the present invention to enhance plant growth, various forms of plant growth enhancement or promotion can be achieved. This can occur as early as when plant growth begins from seeds or later in the life of a plant. For example, plant growth according to the present invention encompasses greater yield, increased quantity of seeds produced, increased percentage of seeds germinated, increased plant size, greater biomass, more and bigger fruit, earlier fruit coloration, and earlier fruit and plant maturation. As a result, the present invention provides significant economic benefit to growers. For example, early germination and early maturation permit crops to be grown in areas where short growing seasons would otherwise preclude their

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growth in that locale. Increased percentage of seed germination results in improved crop stands and more efficient seed use. Greater yield, increased size, and enhanced biomass production allow greater revenue generation from a given plot of land.

Another aspect of the present invention is directed to effecting any form of insect control for plants. For example, insect control according to the present invention encompasses preventing insects from contacting plants to which the hypersensitive response elicitor has been applied, preventing direct insect damage to plants by feeding injury, causing insects to depart from such plants, killing insects proximate to such plants, interfering with insect larval feeding on such plants, preventing insects from colonizing host plants, preventing colonizing insects from releasing phytotoxins, etc. The present invention also prevents subsequent disease damage to plants resulting from insect infection.

The present invention is effective against a wide variety of insects. European com borer is a major pest of com (dent and sweet com) but also feeds on over 200 plant species including green, wax, and lima beans and edible soybeans, peppers, potato, and tomato plus many weed species. Additional insect larval feeding pests which damage a wide variety of vegetable crops include the following: beet armyworm, cabbage looper, com ear worm, fall armyworm, diamondback moth, cabbage root maggot, onion maggot, seed com maggot, pickleworm (melonworm), pepper maggot, and tomato pinworm. Collectively, this group of insect pests represents the most economically important group of pests for vegetable production worldwide.

Another aspect of the present invention is directed to imparting stress resistance to plants. Stress encompasses any environmental factor having an adverse effect on plant physiology and development. Examples of such environmental stress include climate-related stress (e.g., drought, water, frost, cold temperature, high temperature, excessive light, and insufficient light), air polllution stress (e.g., carbon dioxide, carbon monoxide, sulfur dioxide, NO_x, hydrocarbons, ozone, ultraviolet radiation, acidic rain), chemical (e.g., insecticides, fungicides, herbicides, heavy metals), and nutritional stress (e.g., fertilizer, micronutrients, macronutrients). Use of hypersensitive response elicitors in accordance with the present invention impart resistance to plants against such forms of environmental stress.

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The method of the present invention involving application of the hypersensitive response elicitor polypeptide or protein can be carried out through a variety of procedures when all or part of the plant is treated, including leaves, stems, roots, propagules (e.g., cuttings), etc. This may (but need not) involve infiltration of the hypersensitive response elicitor polypeptide or protein into the plant. Suitable application methods include high or low pressure spraying, injection, and leaf abrasion proximate to when elicitor application takes place. When treating plant seeds, in accordance with the application embodiment of the present invention, the hypersensitive response elicitor protein or polypeptide can be applied by low or high pressure spraying, coating, immersion, or injection. Other suitable application procedures can be envisioned by those skilled in the art provided they are able to effect contact of the hypersensitive response elicitor polypeptide or protein with cells of the plant or plant seed. Once treated with the hypersensitive response elicitor of the present invention, the seeds can be planted in natural or artificial soil and cultivated using conventional procedures to produce plants. After plants have been propagated from seeds treated in accordance with the present invention, the plants may be treated with one or more applications of the hypersensitive response elicitor protein or polypeptide to impart disease resistance to plants, to enhance plant growth, to control insects on the plants, and/or impart stress resistance.

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The hypersensitive response elicitor polypeptide or protein can be applied to plants or plant seeds in accordance with the present invention alone or in a mixture with other materials. Alternatively, the hypersensitive response elicitor polypeptide or protein can be applied separately to plants with other materials being applied at different times.

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A composition suitable for treating plants or plant seeds in accordance with the application embodiment of the present invention contains a hypersensitive response elicitor polypeptide or protein in a carrier. Suitable carriers include water, aqueous solutions, slurries, or dry powders. In this embodiment, the composition contains greater than 500 nM hypersensitive response elicitor polypeptide or protein.

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Although not required, this composition may contain additional additives including fertilizer, insecticide, fungicide, nematacide, and mixtures thereof.

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Suitable fertilizers include (NH₄)₂NO₃. An example of a suitable insecticide is Malathion. Useful fungicides include Captan.

Other suitable additives include buffering agents, wetting agents, coating agents, and abrading agents. These materials can be used to facilitate the process of the present invention. In addition, the hypersensitive response elicitor polypeptide or protein can be applied to plant seeds with other conventional seed formulation and treatment materials, including clays and polysaccharides.

In the alternative embodiment of the present invention involving the use of transgenic plants and transgenic seeds, a hypersensitive response elicitor polypeptide or protein need not be applied topically to the plants or seeds. Instead, transgenic plants transformed with a DNA molecule encoding a hypersensitive response elicitor polypeptide or protein are produced according to procedures well known in the art.

The vector described above can be microinjected directly into plant cells by use of micropipettes to transfer mechanically the recombinant DNA.

Crossway, Mol. Gen. Genetics, 202:179-85 (1985), which is hereby incorporated by reference. The genetic material may also be transferred into the plant cell using polyethylene glycol. Krens, et al., Nature, 296:72-74 (1982), which is hereby incorporated by reference.

Another approach to transforming plant cells with a gene which imparts resistance to pathogens is particle bombardment (also known as biolistic transformation) of the host cell. This can be accomplished in one of several ways. The first involves propelling inert or biologically active particles at cells. This technique is disclosed in U.S. Patent Nos. 4,945,050, 5,036,006, and 5,100,792, all to Sanford et al., which are hereby incorporated by reference. Generally, this procedure involves propelling inert or biologically active particles at the cells under conditions effective to penetrate the outer surface of the cell and to be incorporated within the interior thereof. When inert particles are utilized, the vector can be introduced into the cell by coating the particles with the vector containing the heterologous DNA. Alternatively, the target cell can be surrounded by the vector so that the vector is carried into the cell by the wake of the particle. Biologically active particles (e.g.,

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dried bacterial cells containing the vector and heterologous DNA) can also be propelled into plant cells.

Yet another method of introduction is fusion of protoplasts with other entities, either minicells, cells, lysosomes or other fusible lipid-surfaced bodies. Fraley, et al., <u>Proc. Natl. Acad. Sci. USA</u>, 79:1859-63 (1982), which is hereby incorporated by reference.

The DNA molecule may also be introduced into the plant cells by electroporation. Fromm et al., <u>Proc. Natl. Acad. Sci. USA</u>, 82:5824 (1985), which is hereby incorporated by reference. In this technique, plant protoplasts are electroporated in the presence of plasmids containing the expression cassette. Electrical impulses of high field strength reversibly permeabilize biomembranes allowing the introduction of the plasmids. Electroporated plant protoplasts reform the cell wall, divide, and regenerate.

Another method of introducing the DNA molecule into plant cells is to infect a plant cell with Agrobacterium tumefaciens or A. rhizogenes previously transformed with the gene. Under appropriate conditions known in the art, the transformed plant cells are grown to form shoots or roots, and develop further into plants. Generally, this procedure involves inoculating the plant tissue with a suspension of bacteria and incubating the tissue for 48 to 72 hours on regeneration medium without antibiotics at 25-28°C.

Agrobacterium is a representative genus of the gram-negative family Rhizobiaceae. Its species are responsible for crown gall (A. tumefaciens) and hairy root disease (A. rhizogenes). The plant cells in crown gall tumors and hairy roots are induced to produce amino acid derivatives known as opines, which are catabolized only by the bacteria. The bacterial genes responsible for expression of opines are a convenient source of control elements for chimeric expression cassettes. In addition, assaying for the presence of opines can be used to identify transformed tissue.

Heterologous genetic sequences can be introduced into appropriate plant cells, by means of the Ti plasmid of A. tunefaciens or the Ri plasmid of A. rhizogenes. The Ti or Ri plasmid is transmitted to plant cells on infection by Agrobacterium and is stably integrated into the plant genome. J. Schell, Science, 237:1176-83 (1987), which is hereby incorporated by reference.

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After transformation, the transformed plant cells must be regenerated.

Plant regeneration from cultured protoplasts is described in Evans et al., <u>Handbook of Plant Cell Cultures</u>, <u>Vol. 1</u>: (MacMillan Publishing Co., New York, 1983); and Vasil I.R. (ed.), <u>Cell Culture and Somatic Cell Genetics of Plants</u>, Acad. Press, Orlando, Vol. I, 1984, and Vol. III (1986), which are hereby incorporated by reference.

It is known that practically all plants can be regenerated from cultured cells or tissues, including but not limited to, all major species of sugarcane, sugar beets, cotton, fruit trees, and legumes.

Means for regeneration vary from species to species of plants, but generally a suspension of transformed protoplasts or a petri plate containing transformed explants is first provided. Callus tissue is formed and shoots may be induced from callus and subsequently rooted. Alternatively, embryo formation can be induced in the callus tissue. These embryos germinate as natural embryos to form plants. The culture media will generally contain various amino acids and hormones, such as auxin and cytokinins. It is also advantageous to add glutamic acid and proline to the medium, especially for such species as corn and alfalfa. Efficient regeneration will depend on the medium, on the genotype, and on the history of the culture. If these three variables are controlled, then regeneration is usually reproducible and repeatable.

After the expression cassette is stably incorporated in transgenic plants, it can be transferred to other plants by sexual crossing. Any of a number of standard breeding techniques can be used, depending upon the species to be crossed.

Once transgenic plants of this type are produced, the plants themselves can be cultivated in accordance with conventional procedure with the presence of the gene encoding the hypersensitive response elicitor resulting in disease resistance, enhanced plant growth, control of insects on the plant, and/or stress resistance.

Alternatively, transgenic seeds are recovered from the transgenic plants. These seeds can then be planted in the soil and cultivated using conventional procedures to produce transgenic plants. The transgenic plants are propagated from the planted transgenic seeds under conditions effective to impart disease resistance to plants, to enhance plant growth, to control insects, and/or to impart stress resistance. While not

wishing to be bound by theory, such disease resistance, growth enhancement, insect control, and/or stress resistance may be RNA mediated or may result from expression of the elicitor polypeptide or protein.

When transgenic plants and plant seeds are used in accordance with the present invention, they additionally can be treated with the same materials as are used to treat the plants and seeds to which a hypersensitive response elicitor polypeptide or protein is applied. These other materials, including hypersensitive response elicitors, can be applied to the transgenic plants and plant seeds by the above-noted procedures, including high or low pressure spraying, injection, coating, and immersion. Similarly, after plants have been propagated from the transgenic plant seeds, the plants may be treated with one or more applications of the hypersensitive response elicitor to impart disease resistance, enhance growth, control insects, and/or to impart stress resistance. Such plants may also be treated with conventional plant treatment agents (e.g., insecticides, fertilizers, etc.).

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EXAMPLES

Example 1 - Bacterial Strains and Plasmids

20 Escherichia coli DH5 and BL21 were purchased from Gibco BRL (Rockville, MD) and Novagen (Madison, WI) respectively.

pET28 plasmids were from Novagen (Madison, WI).

All restriction enzymes (e.g., NdeI and HindIII), T4 DNA ligase, Calf intestinal alkaline phosphatase (CIP), and PCR reagents were from Gibco BRL (Rockville, MD).

Oligonucleotides were synthesized by Lofstrand Labs Ltd (Gaithersburg, MD).

Chemically synthesized polypeptides were synthesized by Bio-Synthesis (Lewisville, TX).

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Example 2 - Construction of Truncated Gene Encoding Harpin

Fragments of genes encoding harpin proteins were constructed in pET28 vector and expressed in E. coli as follows;

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- HrpN fragments were PCR amplified from the pCPP2139
 plasmid (Cornell University, Ithaca, NY) and cloned into
 pET28 vector.
- HrpZ fragments were PCR amplified from the pSYH10
 plasmid (Cornell University, Ithaca, NY) and cloned into
 pET28 vector.
- PopA fragments were PCR amplified from the pBS::popA
 plasmid (Cornell University, Ithaca, NY) and cloned into
 pBT28 vector.
- HrpW fragments were PCR amplified from the pCPP1233
 plasmid (Cornell University, Ithaca, NY) and cloned into
 pET28 vector.

All truncated fragments were amplified by PCR with full length harpin DNA as the template.

Oligonucleotides corresponding to the truncated N-terminal sequence were started /modified with a Nde I site (which serves as an initiation codon of methionine (ATG)). Oligonucleotides corresponding to a C-terminal sequence contained a UAA stop codon followed by a Hind III site.

PCR was carried in a 0.5 ml tube with GeneAmpTM 9600 and 9700 (PE Applied Biosystems, Branchburg, New Jersey). 45 µl of SuperMixTM (Gibco BRL, Rockville, MD) was mixed with 20 pmoles of each pair of DNA primers, 10 ng of full length harpin DNA, and diH₂O to fill the final volume to 50 µl. After heating the mixture at 95°C for 2 min., PCR was performed for 30 cycles at 94°C for 1 min., 58°C for 1 min. and 72°C for 1.5 min. Amplified DNAs were purified with QIAquick PCR purification kit (QIAGEN Inc., Vlencia, CA), digested with Nde I and Hind III at 37°C for 5 hours, extracted once with phenol:chloroform:isoamylalcohol (25:24:1), and precipitated with ethanol. 5 µg of pET28(b) vector DNA was digested with 15 units of Nde I and 20 units of Hind III at 37°C for 3 hours followed with calf intestinal alkaline phosphatase treatment for 30 min. at 37°C to reduce the background resulting from incomplete single enzyme digestion. Digested vector DNA was purified with the QIAquick PCR purification kit and directly used for ligation. Ligation was carried at 14°C for 12 hours in a 15 µl mixture containing about 50 to

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100 ng of digested pET28(b), 10 to 30 ng of targeted PCR fragments, and 1 unit of T4 DNA ligase. 5 μl of ligation solution was added to 100 μl of DH5α/XL1-Blue competent cells, placed in 15 ml Falcon tube, and incubated on ice for 30 min. After heat shock at 42°C for 45 seconds, 0.9 ml SOC solution (20 g bacto-tryptone, 5 g bacto-yeast extracts, 0.5 g NaCl, 20 mM glucose in one liter) was added into the tube and incubated at 37°C for 1 hour. 20 μl of transformed cells were plated onto LB agar plate with 30 μg/ml of kanamycin and incubated at 37°C for 14 hours. Single colonies were transferred to 3 ml LB-media and incubated overnight at 37°C. Plasmid DNA was prepared in a 2 ml culture with QIAprep Miniprep kit according to the manufacture's instruction. The DNA sequence of truncated harpin constructions was verified with restriction enzyme analysis and sequencing analysis. Plasmids with the desired DNA sequence were transferred into the BL21 strain with a standard chemical transformation method as indicated above.

15 Example 3 - Expression of Proteins

A single clone of *E. coli* with a constructed gene was grown overnight at 37°C in LB with kanamycin. A proper amount of overnight culture was transferred to 50 to 500 ml LB and incubated at 37°C until OD600 reached 0.5 to 0.8. ITPG was added to the culture which was further incubated at room temperature for a period of 5 hour to overnight. Alternatively, a proper amount of overnight culture was transferred to 50 to 500 ml of ½ TB with lactose medium (6 g bacto-trypton, 12 g bacto-yeast extract, 75 g lactose in one liter). After incubation at 37°C until the OD600 reached 0.5 to 0.8, the culture was incubated at room temperature for a period of 5 hours to overnight.

All bacterial cells were harvested by centrifugation and resuspended in 1:5 TE buffer (10 mM Tris, pH 8.5 and 1 mM EDTA). The cells were disrupted by sonication and clarified by centrifugation. Supernatants were then infiltrated into tobacco leaves for HR testing.

Heat treatment (i.e. boiling for 1 to 10 min.) was used to achieve further purification.

All truncated fragments of genes encoding harpin protein were expressed in E. colil BL-21, DE3 strain with an N-terminal His-tag and 20 to 21

amino acid residues generated from the expression vector sequence. The His-tag sequence did not affect the HR activity of the proteins. In some cases, Ni-Agarose beads were added into supernatant solution and mixed at 4°C to room temperature for a period of 30 min. to overnight. The proteins bound to the Ni-Agarose beads were washed by 0.1 M imidazole buffer, and proteins were cluted with 0.6 to 1.0 M imidazole. After dialysis against 10 mM Tris, pH 8.5 buffer, the proteins were infiltrated into tobacco leaves for HR testing.

For proteins expressed in E. coli that were difficult to dissolve in water, total cells were resuspended and sonicated in 8 M urea buffer (0.1M Naphosphate, 10 mM Tris buffer, pH8.0). The total cell lysate was centrifuged, and supernatants were collected. Ni-agarose was added into the supernatants and mixed gently at room temperature for 30 min. The Ni-agarose resin was washed with buffer (8 M urea, 0.1 M Na-phosphate, 10 mM Tris buffer, pH6.3). The proteins were eluted with elution buffer (8 M urea, 0.1 M EDTA, 0.1 M Na-phosphate, 10 mM Tris buffer, pH 6.3) and dialyzed against buffer (pH 8.5, 10 mM Tris) with stepwise decreased urea. If the proteins still were insoluble in buffer, the solution pH was adjusted to 9 to 11 and sonicated at room temperature for 1 to 5 min.

Chemically synthesized polypeptides were dissolved in 10 mM Tris, pH 6.5 to 11 buffers depending on their solubility.

A hypersensitive response ("HR") assay was performed by infiltration of 0.1 to 0.3 ml of serial diluted protein solutions into tobacco leaves (cv. Xanth). All HR data shown in these examples were recorded from 48 hours after infiltration.

Example 4 - Quantification of Proteins

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All expressed proteins were checked with pre-cast 4-20% SDS polyacrylamide gel electrophoresis (SDS-PAGE) from Novex (San Diego, CA). After electrophoresis, the gel was stained with Coomassie R-250 solution (0.1% Coomassie R-250, 10% Acetate Acid, 40% ethanol) for 1 to 4 hours and distained with distaining solution (8% acetate acid and 25% ethanol) overnight. The density of corresponding bands were compared to standard proteins, which were either purchased from Novex or were from quantitative standard harpin protein produced by Eden Bioscience (Bothell, Washington).

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Example 5 - Classification of Harpin Proteins

Since harpin proteins share common biochemical and biophysical characteristics as well as biological functions, based on their unique properties, HR elicitors from various pathogenic bacteria should be viewed as belonging to a new protein family—i.e. the harpin protein family. The harpin protein can be classified into at least four subfamilies based on their primary structure and isolated sources. As set forth in Table 1, those subfamilies are identified by the designation N, W, Z, A,

Table 1 - Subfamilies of Harpin Proteins

Harpin proteins	Isolated Source	Classified Subfamily	pI	Amino acida	Heat stable	Core structure	
77-37	E, amylovora	N	4.42	403	Yes	No	
HrpN _{Ea}	E. chrysanthemi	N	6.51	340	Yes	No	
HrpN _{Bch}	E. carotovora	N	5.82 -	356	Yes	No	
HrpN _{Est}	E. stewartii	N	N/A	N/A	Yes	No	
HrpW _{Ps}	P. syringae	w	4.43	424	Yes	No	
HrpW _{B4}	E. amylovora	w	4.46	447	Yes	No ·	
HrpZ _{Pss}	P. syringae	Z	3.95	341	Yes	No	
PopA1	R.solanacearum	A	4.16	344	Yes	No	

15 <u>Example 6</u> - Analysis of the Structural Units of an HR Domain

The sequence of amino acids that alone could elicit a hypersensitive response in plants (i.e. HR domains) has been investigated in different ways. It was reported that a carboxyl-terminal 148 amino acid portion of HrpZ_{Ps} is sufficient and necessary for HR (He et al., "Pseudomonas Syringae pv. Syringae Harpin_{ps}: A Protein that is Secreted via the Hrp Pathway and Elicits the Hypersensitive Response in Plants," Cell 73:1255-1266.(1993), which is hereby incorporated by reference). With truncated HrpZ fragments, it was determined that an N-terminal 109 amino acids and C-terminal 216 amino acids of HrpZ_{Pss}, respectively, were found to elicit HR (Alfano et al., "Analysis of the Role of the Pseudomonas Syringae pv. Syringae HrpZ Harpin in Elicitation of the Hypersensitive Response in Tobacco Using

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Functionally Non-polar hrpZ Deletion Mutations, Truncated HrpZ Fragments, and hrmA Mutations," Molecular Microbiology 19:715-728 (1996), which is hereby incorporated by reference). Jin et al., "A Truncated Fragment of Harpings Induces Systemic Resistance to Xanthomonas campestris pv. Oryzae in Rice," Physiological and Molecular Plant Pathology 51:243-257 (1997), which is hereby incorporated by reference, reported that a truncated HrpZ_{Ps} with an N-terminal of 137 amino acids elicited a hypersensitive response in tobacco and induced systemic acquired resistance (i.e. SAR) in rice. After digestion with protease, a hypersensitive response active fragment of HrpN_{Es} was isolated and found to span amino acids 137 to 204 of HrpN_{Es}. It was found that a 98 residue of N-terminal HrpN_{Es} fragment was the smallest bacterially produced peptide that displayed HR-eliciting activity (Laby, "Molecular Studies on Interactions Between Erwinia Amylovora and its Host and Non-host Plants," Doctoral Thesis in Cornell University (1997), which is hereby incorporated by reference).

A series of $HrpN_{Ea}$ fragments have been generated with His-tag fusion at the N-terminal of the polypeptides and a polypeptide ($HrpN_{Ea}$ 137180), located at position of 137 to 180 amino acid residue of $HrpN_{Ea}$, was identified to elicit HR activity in tobacco.

20 Example 7 - Analysis of Secondary Structure of HR Domains

The DNA and primary protein sequence of the $HrpN_{Ba}$ 137180 show no any homologues among other hypersensitive response elicitors.

Analyses of the secondary structure of the fragment of $HrpN_{E}$ 137180 revealed, with the aid of the computer program Clone Manger5 (Scientific & Educational Software, Durham, NC), that there was a beta-form, a beta-turn, and unordered forms. One typical α -helical segment of residues at 157-170 was found in the $HrpN_{Ea}$ 137180 polypeptide. To determine the function of this structure, polypeptides with a disrupted α -helical structure were generated and hypersensitive response results were evaluated. As shown in Table 2, a complete alpha-helix unit (H unit), probably with a length greater than 12 amino acid residues, is need for hypersensitive response activity.

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Table 2 - Effect of Alpha-helix Structure

Fragment name HrpN _{Ba} 137180	Amino acid 137-180 (44) pI= 3.10	HR* + <5 μg/ml	Structure Complete H	Source E.coli expressed peptide
HrpN _{Ba} 137166	137-166 (30) pI = 3.29		diarupted H	Synthesized peptide
HrpN _{Ba} 76168	76-168 pI = 3.39	-	disrupted H	E.coli expressed peptide

The α -helical unit plays an important role in hypersensitive response activity; however, it was found that an α -helix unit alone did not achieve HR (Table 3).

Therefore, hypersensitive response eliciting domains contain more than one structure unit. Besides the core α -helical unit, there is an acidic unit that has no typical secondary structure feature but is rich in acidic amino acids. This relaxed structure, having a sheet and random turn, is designated as an acidic unit (A unit).

Although the acidic unit is important in achieving a hypersensitive response, it alone, like the α-helical unit alone, did not elicit a hypersensitive response.

A synthetic polypeptide, HrpN_{Ea}140176, that included both A and H structure, spanning amino acids 140 to 176 of HrpN_{Ea}, gave full activity of HR. Sequence analysis by major search engines revealed no global primary sequence similarity in the databases to HrpN_{Ea}140176, even among the harpin protein families.

Table 3 - Effect of Acidic Unit on Hypersensitive Response (HR) Activity

Fragment name	Amino acid	HR*	Structure (A or H)**	Source
HrpN _{Ea} 140176	140-176 (37) pI=3.17	+ <5 μg/ml	A+H	Synthesized peptide
HrpN _{Es} 157170	157-170 (14) pI = 6.94	-	H	Synthesized peptide
HrpN _{Pa} 137156	137-156 (20) pI = 2.67	-	A	Synthesized peptide

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Example 8 - Hypersensitive Response Domain Structure of HrpNE

Four α -helical regions with at least 12 amino acid residues were found in HrpN_{Ea} based on computer analysis with the program Clone Manager 5 (Scientific & Educational Software, Durham, NC), which predicts the secondary structure of protein from the primary sequence by the method of Garnier-Osguthorpe-Robson.

It is believed that a hypersensitive response domain includes two structural units, the α-helix (H) and the acidic unit (A). Another hypersensitive response domain, spanning amino acids 43 to 70 in HrpNEa, was found. A minimal sequence of 12 to 14 AA residues of both the H and A units is believed to be needed. The chemically synthesized polypeptide of HrpN_{Bs}4370 gave full HR activity in tobacco. Thus, a second HR domain has been discovered based on purely secondary structure analysis and prediction.

To further test the hypothesis that the A and H units are needed to achieve a hypersensitive response, an approach of unit exchange (i.e. swapping an acidic unit from one HR domain to another HR domain) was designed. A polypeptide of HrpN_{Ea}Dswap, which consisted of the acidic unit of a hypersensitive response domain (HrpN_{Ea}140176), spanning amino acids 136 to 156 of HrpN_{Ea}, and the α-helical unit of another hypersensitive response domain (HrpN_{Ea}4370), spanning amino acids 57 to 70 of HrpN_{Ea}, was chemically synthesized. This polypeptide swapped two structural units of A and H between two hypersensitive response domains of HrpN_{Ea}4370 and HrpN_{Ea}140176. The HrpN_{Ea}Dswap gave a hypersensitive response activity in tobacco (Table 4). This result shows that the structural characteristic of an HR domain determines its activity, and structural analysis can be used to determine hypersensitive response activity.

Table 4 - Two Structural Units Determine Hypersensitive Response Activity

Fragment name	Amino acid	HR	Structure Type	Source
НгрN _{в.} 4370	43-70 (28) pI= 3.09	+ <5 µg/ml	A+H	Synthesized peptide Partial soluble
HrpN ₂ ,Dswap	HrpN136156 (A)+ HrpN5770 (H) pI=2.67	<20 µg/ml	A unit from HrpN _{Es} 140176+ H unit from HrpN _{Es} 4370	Synthesized peptide Partial soluble

Example 9 - Prediction of Hypersensitive Response Domains Among Proteins in Harpin Family

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The secondary structure which indicates the presence of a hypersensitive response domain in HrpNEa was used to identify other harpin proteins, including proteins classified as different subfamilies. Structural prediction of a hypersensitive response domain among harpin proteins was carried according to following criteria:

- There are two structural units in a hypersensitive response domain, including:
 - a. A stable α -helix unit with 12 or more amino acids in length and

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- b. An hydrophilic, acidic unit with 12 or more amino acids in length which could be a beta-form, a beta-turn, and unordered forms.
- The pI of a hypersensitive response domain should be acidic and, in general, below 5.

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3. The minimal size of an HR domain is from about 28 to 40 AA residues.

Putative HR domains have been identified to fit the criteria by computer analysis among harpin protein family (Table 5).

Table 5 - Predication of Hypersensitive Response Domains Among Harpin
Proteins

HR domain	Isolated Source	Predicted region*	pI	Structure
HroN _B -1	E. amylovora	43-70	3.09	A+H
HrpN _{Ea} -2	E. amylovora	140-176	3.17	A+H
HrpNpa-1	E. chrysanthemi	78-118	5.25	A+H
HrpN _{Ech} -2	E. chrysanthemi	256-295	4.62	A+H
HrpN _{Ecc} -1	E. carotovora	25-63	4.06	A+H
HrpN _{Boe} -2	E. carotovora	101-140	3.00	A+H
		52-96	4.32	A+H
HrpW _{Pss} -1	P. syringae	10-59	4.53	A+H
HrpW _{Pa} -1	E. amylovora	10-39	1	
HrpZ _{Pss} -1	P. syringae	97-132	3.68	A+H
HrpZ _{Pu} -2	P. syringae	153-189	3.67	A+H
HrpZ _{Pss} -3	P. syringae	271-308	3.95	A+H
				- 1 TT
PopAl _{Re-1}	R.solanacearum	92-125	3.75	A+H
PopAl _{Re-2}	R.solanacearum	206-260	3.62	A+H_

Amino acid residue position

Example 10 - Hypersensitive Response Activity of Select Synthesized Polypeptides

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Polypeptides were produced by expression in either E. coli or by chemical synthesis. Based on prediction of solubility and stability of a particular peptide, in some cases, a broader region of AA residues in addition to the essential units were also synthesized to increase solubility of the peptides. The identification of HR domains among four subfamilies of harpin protein demonstrated this (Table 6).

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Table 6 - Hypersensitive Response Activity of Select Synthesized Polypeptides

HR domain	Isolated Source	Synthesized region	pI	Source	HR activity
HrpNg-1	E. amylovora	43-70	3.09	Chemical Synthesized	+ < 5 μg/ml
HrpN _{Bs} -2	E. amylovora	140-176	3.17	Chemical Synthesized	+<5 μg/ml
HrpW _{Es} -2	E. amylovora	10-59	4.53	E.coli expressed	+ < 5 μg/ml
HrpZ _{Pss} -1	P. syringae	97-132	3.68	Chemical Synthesized	+ < 20 µg/ml
HrpZ _{Psr} -1	P. syringae	153-189	3.69	E.coli expressed	+ < 5 μg/ml
PopAl _{Rs} -1	R. solanacearum	92-125	3.75	Chemical Synthesized	+ < 5 μg/ml
PopA1 _{Rs} -2	R. solanacearim	206-260	3.62	E.coli expressed	+<5 µg/ml

5 Example 11 - Construction of Hypersensitive Response Domains in a Protein Expression Cassette

Polypeptides with a harpin protein hypersensitive response domain were expressed in E. coli. PCR was used to amplify desired areas of genes encoding harpin proteins and cloned into an expression vector, e.g. pET28a. A pair of PCR primers with unique flanking sequences were designed to create a universal expression cassette, as shown in Figure 1, for expression of a fragment of harpin protein. Each amplified DNA fragment has a protein translation start codon of ATG in a restriction enzyme Nde I site which might add an extra amino acid of methionine into a polypeptide. Each amplified DNA fragment has a protein translation stop codon of TAA. Each amplified fragment contained two restriction enzyme sites of EcoR V and Sma I, which gave 4 extra in-frame amino acids expressed as Pro-Gly at the N-terminal and Asp-Ile at the C-terminal, respectively. Those two sites are essential to allow two or more expression cassettes to be linked in a specific order and in frame with a minimum number of amino acids being introduced. Cassette A was first digested by EcoR V, ligated to cassette B, and digested with Sma I to produce a new expression cassette C which coupled the two fragments together with two extra amino acids (i.e. Asp-Gly), which are common amino acids in hypersensitive response domains. The newly formed cassette C still contained the same 5' and 3' flanking sequences as original cassettes A and B and maintained the ability to be

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coupled by another cassette. Bgl II and Bam HI sites in the cassette permit the cassette to be linked in frame into a cancatomer with a correct orientation. The strategy is that digestion of DNA with Bgl II and Bam HI results in compatible ends that would be ligated with each other but could not be cut by either enzymes after ligation. For example, a DNA fragment encoding a hypersensitive response domain in a cassette could be digested by restrictions enzymes of Bgl II and Bam H1 separately, digested DNA fragments could be ligated in a ligation solution also including both Bgl II and Bam HI enzymes, any ligated ends with Bgl II or Bam HI sites could be digested by the enzymes, and only those ligated sites between Bgl II and Bam HI could remain.

Example 12 - Building Blocks for Creating Superharpins that have Higher Biological Efficacy

Hypersensitive response domains were identified and isolated from several harpin proteins. With the combination of those HR domains, new polypeptides (i.e. superharpins) that have higher HR potency and have enhanced ability to induce disease resistance, impart insect resistance, enhance growth, and achieve environmental stress tolerance. Superharpins could be one HR domain repeat units (cancatomer), different combinations of HR domains, and/or biologically active domains from other elicitors. Part of the domains from different harpin proteins and other elicitors were constructed into the universal expression cassette as shown on Example 11 and designated as superharpin building blocks. Table 7 lists some superharpin building blocks which were expressed in pET-28a(+) vector with a His-tag sequence at their N-terminal.

Table 7 - Superharpin Building Blocks including pET-28a(+) his-tag Leader Sequence

Domain	Source	MW (kDa)	#a.a.	pI	Soluble	(Structurally) Heat Stable
Sequence	2 150 146	10.69	104	6.48	Yes	Yes
A	PopA70-146	6.754	68	6.78	N/A	N/A
(N _N)	HrpNEa40-80	10.84	111	6.13	N/A	N/A
$(N_N)_2$	Dimer of HrpNEa40-80		154	5.63	N/A	N/A
(N _N) ₃	Triplemer of HrpNEa40-80	14.93		4.95	N/A	N/A
(N _N)4	Tetramer of HrpNEa40-80	19.01	197		Yes	Yes
(N _c)	HrpNEa140-180	7.224	68	5.01		Yes
(N _C) ₂	Dimer of HrpNEa140-180	11.78	111	3.98	Yes	
	Triplemer of HrpNBa140-	16.34	154	3.72	Yes	Yes
(Nc)s	180					
07.	Tetramer of HrpNEa140-	20.89	197	3.58	Yes	Yes
(Nc)+	180	1	1	İ		<u> </u>
	Cancatomer (10 repeating	48.23	455	3.28	N/A	N/A
(Nc)10	units of HrpNEa140-180		1	_		
	Cancatomer (16 repeating	75.57	713	3.18	NA	NVA
(Nc)16	units of HrpNEa140-180				1	İ
			77	6.48	N/A	N/A
W	HrpWEa10-59		78	5.38		Yes
Z _N	HrpZ90-150			6.40		Yes
Z266-308	HrpZ266-308					
his-tag leader		2.045	1 13	11.04		
seq.				ــــــــــــــــــــــــــــــــــــــ		

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<u>Example 13</u> - Superharpins with Stacked HR Domains and their Biological Activities

There are numerous polypeptides could be generated with different

combinations of HR domains or by stacking HR domains and repeating units in order.

Selective combination or stacking of HR domains isolated from harpin proteins or
other elicitors can be designed to achieve a targeted disease resistance spectrum. See
Table 8 for superharpins prepared by stacking of HR building blocks listed on
Table 7. All three listed superharpins (i.e. SH-1, SH-2, SH-3) were constructed into a

pET28(a) vector and expressed in E. coli. Recombinant proteins were partially purified and quantified by SDS-PAGE with purified Harpin N protein as a quantitative standard.

Table 8 - Properties of Superharpins

Protein	Domain Sequence	MW (kDa)	# a.a.	pI	Soluble	Heat Stable
SH-1	*W(N _N)4A(N _C)4Z ₂₆₆₋₃₀₈	54,955	545	3.69	Yes	Yes
SH-2	*W(N _N) ₄ Z _N (N _C) ₄ Z ₂₆₆₋₃₀₈	52.341	519	3.54	Yes	Yes
SH-3	*W(N _N) ₄ Z _N (N _C) ₄ Z ₂₆₆₋₃₀₈ A	60.375	598	3.67	Yes	Yes
HroNEa	HrpN from E.amylovora	39.697	403	4.42	Yes	Yes

Bioassays for hypersensitive response on tobacco leaves (HR), percentage of TMV reduction on tobacco leaves, and plant growth enhancement with tomato showed that superharpins had higher (up to 2 to 10 fold greater) HR potency compared with HrpN from *E. amylovora*. This also demonstrated that superharpins have better performance on % TMV reduction and plant growth enhancement assay. See Table 9.

Table 9 - Biological Activities of Superharpins

Protein	Domain Sequence	Elicit HR	% TMV reduct	on on tobacco	% Plant Growth Enhancement		
Protein	Donain ovq	(~µg/ml) 10 µg/ml		i μg/ml	10 μ g/ml	1 μg/ml	
	77013 (01) 7	0.66	B3	79	7.49	9.83	
SH-1	W(N _N)4A(N _C)4Z266.508	0.13	84	60	11.05	7.30	
SH-2	W(N _N) ₄ Z _M (N _C) ₄ Z ₂₆₆₋₃₆₈	0.15	77	55	11.07	10.00	
SH-3 HmNEa	W(N _N) ₄ Z _N (N _C) ₄ Z ₂₆₆₋₅₀₈ A HrpN from E.amylovora	1-3	55	10	11.68	NA	

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Although the invention has been described in detail for the purpose of illustration, it is understood that such detail is solely for that purpose, and variations can be made therein by those skilled in the art without departing from the spirit and scope of the invention which is defined by the following claims.

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WHAT IS CLAIMED:

- An isolated hypersensitive response elicitor protein comprising an isolated pair or more of spaced apart domains, each comprising an acidic portion linked to an alpha-helix and capable of eliciting a hypersensitive response in plants.
 - 2. A protein according to claim 1, wherein the protein is recombinant.
- 10 3. An isolated nucleic acid molecule encoding a protein according to claim 1.
 - 4. A nucleic acid molecule according to claim 3, wherein each domain is from a different source organism.
 - A nucleic acid molecule according to claim 3, wherein there are
 or more spaced apart domains.
- 20 according to claim 3 which is heterologous to the expression vector.
 - 7. An expression vector according to claim 6, wherein the nucleic acid molecule is positioned in the expression vector in sense orientation and correct reading frame.
 - 8. A host cell transformed with the nucleic acid molecule according to claim 3.
- A host cell transformed according to claim 8, wherein the host
 cell is selected from the group consisting of a plant cell, a eukaryotic cell, and a procaryotic cell.

- 10. A host cell according to claim 8, wherein the nucleic acid molecule is transformed with an expression system.
- 11. A transgenic plant transformed with the nucleic acid molecule of claim 3.
 - 12. A transgenic plant according to claim 11, wherein the plant is selected from the group consisting of alfalfa, rice, wheat, barley, rye, cotton, sumflower, peanut, corn, potato, sweet potato, bean pea, chicory, lettuce, endive, cabbage, brussel sprout, beet, parsnip, cauliflower, broccoli, turnip, radish, spinach, onion, garlic, eggplant, pepper, celery, carrot, squash, pumpkin, zucchini, cucumber, apple, pear, melon, citrus, strawberry, grape, raspberry, pineapple, soybean, tobacco, tomato; sorghum, and sugarcane.
- 15 13. A transgenic plant according to claim 11, wherein the plant is selected from the group consisting of Arabidopsis thaliana, Saintpaulia, petunia, pelargonium, poinsettia, chrysanthemum, camation, and zinnia.
- 14. A transgenic plant according to claim 11, wherein the plant is a20 monocot.
 - 15. A transgenic plant according to claim 11, wherein the plant is a dicot.
- 25 16. A transgenic plant according to claim 11, wherein each domain is from a different source organism.
 - 17. A transgenic plant according to claim 11, wherein there are 3 or more spaced apart domains.
 - 18. A transgenic plant seed transformed with the nucleic acidemolecule of claim 3.

- 19. A transgenic plant seed according to claim 18, wherein the plant is selected from the group consisting of alfalfa, rice, wheat, barley, rye, cotton, sunflower, peanut, corn, potato, sweet potato, bean pea, chicory, lettuce, endive, cabbage, brussel sprout, beet, parsnip, cauliflower, broccoli, turnip, radish, spinach, onion, garlic, eggplant, pepper, celery, carrot, squash, pumpkin, zucchini, cucumber, apple, pear, melon, citrus, strawberry, grape, raspberry, pineapple, soybean, tobacco, tomato, sorghum, and sugarcane.
- 20. A transgenic plant seed according to claim 18, wherein the plant is selected from the group consisting of Arabidopsis thaliana, Saintpaulia, petunia, pelargonium, poinsettia, chrysanthemum, carnation, and zinnia.
- 21. A transgenic plant seed according to claim 18, wherein the plant is a monocot.
 - 22. A transgenic plant seed according to claim 18, wherein the plant is a dicot.
- 23. A method of imparting disease resistance to plants comprising: applying a protein according to claim 1 to a plant or a plant seed under conditions effective to impart disease resistance to the plant or to a plant grown from the plant seed.
- 25 24. A method according to claim 23, wherein the protein is applied to a plant.
 - 25. A method according to claim 23, wherein the protein is applied to a plant seed and further comprising:
- planting the plant seed under conditions effective to impart disease resistance to a plant grown from the plant seeds.

		A method of enhancing plant growth comprising:
	applyi	ng a protein according to claim 1 to a plant or a plant seed under
conditions eff	ective to	enhance growth of the plants or of a plant grown from the plant
seed.		

- 27. A method according to claim 26, wherein the protein is applied to a plant.
- 28. A method according to claim 26, wherein the protein is applied to a plant seed and further comprising:

 planting the plant seeds under conditions effective to enhance growth

of a plant grown from the plant seed.

- 29. A method of controlling insects comprising: applying a protein according to claim 1 to a plant or a plant seed under conditions effective to control insects.
- 30. A method according to claim 29, wherein the protein is applied to a plant.

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31. A method according to claim 29, wherein the protein is applied to a plant seed and further comprising:

planting the plant seed under conditions effective to grow a plant from the plant seed and to control insects.

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32. A method of imparting stress resistance to plants comprising: applying a protein according to claim 1 to a plant or a plant seed under conditions effective to impart stress resistance to the plant or to a plant grown from the plant seed.

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33. A method according to claim 32, wherein the protein is applied to a plant.

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- 34. A method according to claim 32, wherein the protein is applied to a plant seed and further comprising:
- planting the plant seed under conditions effective to impart stress

 resistance to a plant grown from the plant seed.
 - 35. A method of imparting disease resistance to plants comprising: providing a transgenic plant or transgenic plant seed containing the nucleic acid according to claim 3 and
 - planting the transgenic plant or transgenic plant seed under conditions effective to impart disease resistance to the plant or to a plant grown from the plant seed.
- 36. A method according to claim 35, wherein a transgenic plant is provided.
 - 37. A method according to claim 35, wherein a transgenic plant seed is provided.
- 20 38. A method of enhancing growth of plants comprising:

 providing a transgenic plant or transgenic plant seed containing the
 nucleic acid according to claim 3 and

planting the transgenic plant or transgenic plant seed under conditions effective to enhance growth of the plant or of a plant grown from the plant seed.

- 39. A method according to claim 38, wherein a transgenic plant is provided.
- 40. A method according to claim 38, wherein a transgenic plant30 seed is provided.
 - 41. A method of controlling insects comprising:

providing a transgenic plant or transgenic plant seed containing the nucleic acid according to claim 3 and

planting the transgenic plant or transgenic plant seed under conditions effective to control insects on the plant or on a plant grown from the plant seed.

- 42. A method according to claim 41, wherein a transgernic plant is provided.
- 43. A method according to claim 41, wherein a transgenic plant 10 seed is provided.
 - 44. A method of imparting stress resistance to plants comprising: providing a transgenic plant or transgenic plant seed containing the nucleic acid according to claim 3 and
- planting the transgenic plant or transgenic plant seed under conditions effective to impart stress resistance to the plant or to a plant grown from the plant seed.
- 45. A method according to claim 44, wherein a transgenic plant is provided.
 - 46. A method according to claim 44, wherein a transgenic plant seed is provided.
- 25 47. An isolated hypersensitive response elicitor protein comprising, in isolation, a domain comprising an acid portion linked to an alpha-helix and capable of eliciting a hypersensitive response in plants.
- 48. A protein according to claim 47, wherein the protein is 30 recombinant.

- 49. An isolated nucleic acid molecule encoding a protein according to claim 47.
- 50. An isolated nucleic acid molecule according to claim 49,
 5 wherein there are at least 2 domains, each from a different source organism.
 - 51. An isolated nucleic acid molecule according to claim 49, wherein there are 3 or more coupled domains.
- 10 52. An expression vector containing a nucleic acid molecule according to claim 49 which is heterologous to the expression vector.
 - 53. An expression vector according to claim 52, wherein the nucleic acid molecule is positioned in the expression vector in sense orientation and correct reading frame.
 - 54. A host cell transformed with the nucleic acid molecule according to claim 49.
- 20 55. A host cell transformed according to claim 54, wherein the host cell is selected from the group consisting of a plant cell, a eukaryotic cell, and a prokaryotic cell.
- A host cell according to claim 54, wherein the nucleic acid
 molecule is transformed with an expression system.
 - 57. A transgenic plant transformed with the nucleic acid molecule of claim 49.
- 30 58. A transgenic plant according to claim 57, wherein the plant is selected from the group consisting of alfalfa, rice, wheat, barley, rye, cotton, sunflower, peanut, corn, potato, sweet potato, bean pea, chicory, lettuce, endive,

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cabbage, brussel sprout, beet, parsnip, cauliflower, broccoli, turnip, radish, spinach, onion, garlic, eggplant, pepper, celery, carrot, squash, pumpkin, zucchini, cucumber, apple, pear, melon, citrus, strawberry, grape, raspberry, pineapple, soybean, tobacco, tomato, sorghum, and sugarcane.

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- 59. A transgenic plant according to claim 57, wherein the plant is selected from the group consisting of *Arabidopsis thaliana*, *Saintpaulia*, petunia, pelargonium, poinsettia, chrysanthemum, carnation, and zinnia.
- 10 60. A transgenic plant according to claim 57, wherein the plant is a monocot.
 - 61. A transgenic plant according to claim 57, wherein the plant is a dicot.

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- 62. A transgenic plant according to claim 57, wherein there are at least 2 coupled domains, each from a different source organism.
- 63. A transgenic plant according to claim 57, wherein there are 3 or more coupled domains.
 - 64. A transgenic plant seed transformed with the nucleic acid molecule of claim 49.
- 25 A transgenic plant seed according to claim 64, wherein the plant is selected from the group consisting of alfalfa, rice, wheat, barley, rye, cotton, sunflower, peanut, corn, potato, sweet potato, bean pea, chicory, lettuce, endive, cabbage, brussel sprout, beet, parsnip, cauliflower, broccoli, turnip, radish, spinach, onion, garlic, eggplant, pepper, celery, carrot, squash, pumpkin, zucchini, cucumber, apple, pear, melon, citrus, strawberry, grape, raspberry, pineapple, soybean, tobacco, tomato, sorghum, and sugarcane.

- 66. A transgenic plant seed according to claim 64, wherein the plant is selected from the group consisting of Arabidopsis thaliana, Saintpaulia, petunia, pelargonium, poinsettia, chrysanthemum, carnation, and zinnia.
- 5 67. A transgenic plant seed according to claim 64, wherein the plant is a monocot.
 - 68. A transgenic plant seed according to claim 64, wherein the plant is a dicot.
- 69. A method of imparting disease resistance to plants comprising: applying a protein according to claim 47 to a plant or a plant seed under conditions effective to impart disease resistance to the plant or to a plant grown from the plant seed.
 - 70. A method according to claim 69, wherein the protein is applied to a plant.
- 71. A method according to claim 69, wherein the protein is applied
 20 to a plant seed and further comprising:

 planting the plant seed under conditions effective to impart disease
 resistance to a plant grown from the plant seed.
- 72. A method of enhancing plant growth comprising:

 applying a protein according to claim 47 to a plant or a plant seed under conditions effective to enhance growth of the plant or of a plant grown from the plant seed.
- 73. A method according to claim 72, wherein the protein is applied 30 to a plant.

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74. A method according to claim 72, wherein the protein is applied to a plant seed and further comprising:

planting the plant seed under conditions effective to enhance growth of a plant grown from the plant seed.

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- 75. A method of controlling insects comprising: applying a protein according to claim 47 to a plant or a plant seed under conditions effective to control insects.
- 76. A method according to claim 75, wherein the protein is appliedto a plant.
 - 77. A method according to claim 75, wherein the protein is applied to a plant seed and further comprising:

planting the plant seed under conditions effective to grow a plant from the plant seed and to control insects.

- 78. A method of imparting stress resistance to plants comprising:
 applying a protein according to claim 47 to a plant or a plant seed
 under conditions effective to impart stress resistance to the plant or to a plant grown from the plant seed.
 - 79. A method according to claim 78, wherein the protein is applied to a plant.

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80. A method according to claim 78, wherein the protein is applied to a plant seed and further comprising:

planting the plant seed under conditions effective to impart stress resistance to a plant grown from the plant seed.

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81. A method of imparting disease resistance to plants comprising:

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providing a transgenic plant or transgenic plant seed containing the nucleic acid according to claim 49 and

planting the transgenic plant or transgenic plant seed under conditions effective to impart disease resistance to the plant or to a plant grown from the plant seed.

- 82. A method according to claim 81, wherein a transgenic plant is provided.
- 10 83. A method according to claim 81, wherein a transgenic plant seed is provided.
- 84. A method of enhancing growth of plants comprising:

 providing a transgenic plant or transgenic plant seed containing the

 nucleic acid according to claim 49 and

 planting the transgenic plant or transgenic plant seed under conditions

 effective to enhance growth of the plant or of a plant grown from the plant seed.
- 85. A method according to claim 84, wherein a transgenic plant is 20 provided.
 - 86. A method according to claim 84, wherein a transgenic plant seed is provided.
- 25 87. A method of controlling insects comprising:

 providing a transgenic plant or transgenic plant seed containing the
 nucleic acid according to claim 49 and
 planting the transgenic plant or transgenic plant seed under conditions

planting the transgenic plant or transgenic plant seed under conditions effective to control insects on the plant or on a plant grown from the plant seed.

88. A method according to claim 87, wherein a transgenic plant is provided.

- 89. A method according to claim 87, wherein a transgenic plant seed is provided.
- 90. A method of imparting stress resistance to plants comprising: providing a transgenic plant or transgenic plant seed containing the nucleic acid according to claim 49 and

planting the transgenic plant or transgenic plant seed under conditions effective to impart stress resistance to the plant or to a plant grown from the plant 10 seed.

- 91. A method according to claim 90, wherein a transgenic plant is provided.
- 92. A method according to claim 90, wherein a transgenic plant seed is provided.

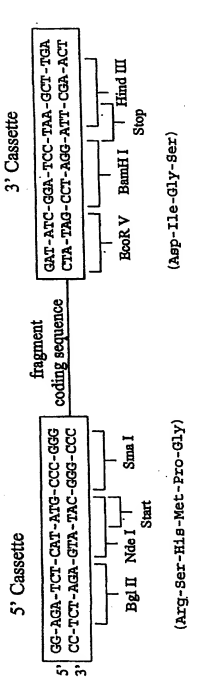


Figure 1

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Leu Leu Gly His Asp Thr Val Thr Lys Leu Thr Asn Gln Ser Asn Gln 115 120 125

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WO 01/98501 PCT/US01/18820

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WO 01/98501

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TOW		

1545

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1800

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1035

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HARPIN FROM ERWINIA AMYLOVORA INDUCES PLANT RESISTANCE

Z.-M. Wei and S. V. Beer Department of Plant Pathology Cornell University Ithaca, NY 14853 USA

Plants have evolved a complex array of biochemical pathways that enable them to recognize and respond to signals from the environment. A common form of plant resistance is the restriction of pathogen proliferation to a small zone surrounding the site of infection. Typically this restriction is accompanied by localized necrosis. In addition to local defense response, plants also respond to infection by activating defenses in uninfected parts of the plant, which result in resistance of the plant to secondary infection (Dean and Kuc, 1985). Collectively, this phenomenon of induced resistance is called systemic acquired resistance (SAR). SAR reduces the severity of disease caused by all classes of pathogens and it can persist for several weeks or longer. SAR can be induced by abiotic agents, such as salicylic acid as well as biotic agents, such as virulent and avirulent pathogens (Dean and Kuc, 1985; Malamy et al., 1990). Salicylic acid is believed to play a signal function in the induction of SAR since endogenous levels of salicylic acid increase after "Immunization" with an incompatible pathogen. However at present, little is known about the signal transduction pathways activated during responses of a plant to attack by a pathogen, although this knowledge is central to understanding disease susceptibility and resistance.

Erwinia amylovora is an often devastating plant pathogenic bacterium that causes the fire blight disease of pear, apple and many other rosaceous plants. In non-host plants, E. amylovora elicits the hypersensitive response (HR), which is characterized by a rapid, localized death of tissues infiltrated with high concentrations of bacterial cells (>10⁷ cfu/ml) (Klement, 1982). hrp genes are essential for E. amylovora to cause disease in host plants and to elicit the HR in non-host plants (Beer et al., 1991). Harpin is a heat-stable, glycine-rich, secreted protein with molecular mass of 37 kD. It is encoded by hrpN of E. amylovora (Wei et al., 1992). When infiltrated into intercellular spaces, harpin elicits the HR in many plants including tobacco, pepper, sunflower, tomato cabbage, arabidopsis, cucumber, geranium, watermelon and lettuce.

The HR is believed to be associated with plant defense against pathogens. Hence, we reasoned that harpin-induced HR may induce plant resistance. We tested harpin-induced resistance in more than seven different plants against eight diseases caused by fungi, bacteria and viruses. All tested plants showed some resistance. Here we report evidence of harpin-induced resistance to three diseases, southern bacterial wilt of tomato, tobacco mosaic virus and Gliocladium leaf spot of cucumber.

Harpin-induced resistance in tomato against southern bacterial wilt caused by Pseudomonas solanacearum.

100 μ l of a cell suspension of ca. 108 ctu/ml of Escherichia coli DH5 α (pCPP430) or 100 μ l of a 200 μ g/ml crude harpin preparations were infiltrated into portions of the two lower true leaves of two-week-old tomato seedlings grown in 8 x 15 cm flats in the greenhouse. Twenty plants were used for each treatment. Necrosis was evident 24 hours after infiltration of harpin or *E. coli* DH5 α (pCPP430), which produces and secretes

harpin. Four days after the tomato seedlings had been treated with harpin or bacteria, they were inoculated with P. solanacearum K60 (107 cfu/ml) by root dipping for three minutes. The inoculated plants were replanted into the same flats and left in a greenhouse. None of the 20 harpin-infiltrated plants showed any symptoms one week after inoculation with P. solanacearum K60. However, seven of the 20 buffer-infiltrated plants were stunted. After two weeks, 11 buffer-infiltrated plants showed severe wilting and five were stunted, characteristics of the southern bacterial wilt disease. In comparison, only two harpin-treated plants appeared wilted and three plants were stunted. Similar induced resistance was observed following infiltration of living bacteria E. coli/DH5α(pCPP430), but not by E. coli DH5α(pCPP430), which is a harpin-deficient mutant created by transposon Tn5tac insertion into the hrpN gene. These results indicate that harpin, which is produced and secreted by hrp gene cluster of E. amylovora, is responsible for the induced-resistance realized.

Harpin-induced resistance in tobacco to tobacco mosaic virus (TMV)

One panel of a lower leaf of four-week-old tobacco seedlings (Nicotiana tabacum L. "Xanthi" with N gene) was infiltrated with 100 µl of a 200 µg/ml crude harpin preparation in 5 mM phosphate buffer. Three days later, the plants were challenged with TMV. Fifty µl of a suspension of TMV (5 µg/ml) was rubbed on one upper leaf with 400-mesh carborundum. Six plants were used for each treatment. Necrotic lesions appeared on inoculated leaves of both harpin- and buffer-treated plants 4 days after inoculation. The average number of necrotic lesions from the six harpin-treated plants was 21, which was significantly less than the 67 lesion average that developed on six buffer-treated plants. More importantly, the size of the lesions on buffer-treated plants was larger than those on the harpin-treated plants. Actually, it was difficult to distinguish individual lesions on the buffer-treated plants by day 10, because several necrotic lesions had merged.

Harpin-induced resistance against Gliocladium leaf spot of cucumber

Harpin or a cell suspension of E. coli DH5α(pCPP430) was infiltrated into first two true leaves of two-week-old cucumber seedlings. Six plants were infiltrated for each treatment. Four days after infiltration of harpin, a Gliocladium cucurbitae spore suspension (108 spores/ml) was sprayed onto the whole plants. The inoculated plants were incubated in a moisture chamber. Ten days after the inoculation, typical leaf spots appeared. A mean of six lesions was present on the lowest leaves of six harpin-treated plants, but 32 lesions formed on the same leaves of the six buffer-treated plants. On the third lowest leaves, the difference in disease severity was even greater; there were virtually no lesions on harpin-treated plants, however, more than 30 lesions were found on the buffer-treated plants. Later, most of the diseased leaves on buffer-treated plants

The examples outlined above show that harpin is able to induce resistance in different plants against bacterial, viral and fungal pathogens. Although mechanisms of harpin-induced resistance are unknown, some of our preliminary experiments have shown that harpin may act as an elicitor of salicylic acid induction, which is believed to be involved in SAR (Malamy et al., 1990). Unlike some host-specific elicitors (Keen et al., 1990), harpin is able to elicit the HR on a broad range of plants. Thus, we expect that harpin-induced resistance can be achieved in many plants either by manipulation of harpin exogenously or by harpin-mediated transgenic plants.

Our studies of harpin-induced resistance are just beginning and we need to learn more to understand the exciting features of this phenomenom. For example, what is the minimal amount of harpin needed to induce plant resistance and how long does the resistance persist, and what mechanisms are involved in harpin-induced resistance? We expect that harpin as a novel molecule will play an important role in dissecting the signal transduction pathways of induced-resistance in plants, and perhaps also in practical disease control.

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Induction of systemic acquired resistance in cucumber by Pseudomonas syringae pv. syringae 61 HrpZ_{Pss} protein

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Summerv

Systemic acquired resistance (SAR) is an inducible plant defense response and is effective against a broad spectrum of pathogens. Biological Induction of SAR usually follows plant cell death resulting from the plant hypersonstive response (HR) elicited by an avirulent pathogen or from disease necrosis caused by a virulent pathogen. The elicitation of the HR and disease necroses by pathogenic becteria is controlled by hip genes. Previously, it was shown that the Pseudomonas syringae 61 (Pss61) HrpZ_{Pss} protein (formally harpings) elicited the HR in plants. in this study, it is shown that HrpZ_{PM} induced SAR in eucumber to diverse pathogens, including the anthracnose fungus (Colletotrichum lagenarium), tobacco necrosis virus and the bacterial angular leaf spot bacterium (P. s. pv. lachrymans). A hupH mutant of Pss61, which is defective in the secretion of $HrpZ_{Pas}$ and, possibly, other protein elicitors, failed to elicit SAR. Pathogenesis-related (PR) proteins, including peroxidase, β-glucanase and chitinases, were induced in cucumber plants inoculated with Pss61, C. lagenarium or $HrpZ_{Pos}$. The induction patterns of PR proteins by HrpZ_{Pss} and Pss61 were the same, but were different from that induced by C. legenarium. Interestingly, the hipH mutant induced two of the three identified PR proteins, despite its failure to induce SAR. These results suggest that proteinaceous elicitors, such as HrpZ_{Pas}, that traverse the bacterial Hrp secretion pathway are involved in the biological induction of SAR and that at least some PR proteins can be induced by bacterial factors that are not controlled by hrp genes.

Introduction

Localized infection of plants by necrotizing pathogens can result in systemic acquired resistance (SAR) to disease, which persists for weeks to months and is effective against diverse pathogens including fungi, bacteria, and necrotiz-

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ing viruses (Kuc, 1982; Ross, 1961). Biological induction of SAR is usually associated with prior plant cell death during the hypersensitive response (HR) or disease necrosis triggered by avirulent or virulent pathogens, respectively (Cameron et al., 1994; Kuc, 1982; Ross, 1961; Uknes et al., 1993). Certain synthetic chemicals, such as salicylic add (SA) and 2,6-dichloroisonicotinic acid (INA), also can be very effective in the induction of SAR when applied to plants (Metraux et al., 1991; White, 1979). The induction of SAR in cucumber plants by an avirulent bacterial pathogen, Pseudomonas syringae pv. syringae, appears to be dependent on bacterial hrp genes that are required for many plant pathogenic bacteria to elicit the HR in non-host plants or to cause disease in host plants (Smith et al., 1991). The HR is a complex plant resistance reaction which involves local plant cell death and restriction of pathogens to the site of their introduction (Klement, 1982).

Recent studies have shown that most Hrp proteins are involved in the assembly of a type ill protein secretion pathway (the Hrp pathway) through which bacterial pathogenesis-related proteins traverse to the extracellular milieu to initiate various plant-bacterial interactions (Fenselau, 1992; Huang et al., 1992, 1995; Van Gijsegem et al., 1995). One family of such proteins that have been identified are heat-stable, glycine-rich proteins: harpin of Erwinia amylovora (Wei et al., 1992), HrpZpes (formally harpinges) of P. s. pv. syringae 61 (Pss61) (He et al., 1993) and PopA of P. solanscearum (Ariat et al., 1994). Harpins and PopA were shown to elicit the HR when infiltrated into the leaf laminae of appropriate plants (Arlat et al., 1994; He et al., 1993; Wei et al., 1992), to induce exchange of H+ and K+ (the 'XR') across the plasmalermma (Wei et al., 1992), and to generate active oxygen species (Baker et al., 1993) when added to plant cell cultures, which are all properties of the HR elicited by live bacteria.

As part of our investigation into plant responses to *P. syringae* extracellular proteins under the control of the Hrp regulatory/secretion system, we studied the involvement of HrpZ_{Pss} In the biological induction of SAR by *P. s. pv. syringae* 61. In this paper we describe the experimental results showing that HrpZ_{Pss}, as well as the bacterium (Pss61) that produces it, efficiently induced SAR in cucumber to diverse pathogens, including a fungus (*Colletotrichum lagenarium*), a bacterium (*P. s. pv. lachrymans*) and a local lesion-forming virus (tobacco necrosis virus). The 'hrpH mutant, which is defective in the secretion of HrpZ_{Pss}, failed to induce SAR. Multiple pathogenesis-related (PR) proteins were detected in cucumber plants treated with HrpZ_{Pss}, Pss61 and *C. legenarium*. The efficacy

of SAR induction, resistance spectrum and patterns of PR protein induction were very similar in plants treated with HrpZ_{Pss} and Pss61. Interestingly, the PR protein patterns induced by HrpZ_{Pss} and Pss61 were somewhat different from that induced by C. lagenarium. The hrpH mutant, though unable to induce SAR, efficiently induced some of the well-characterized PR proteins. These results suggest that the biological induction of SAR by P. syringae is dependent on the bacterial proteins (such as HrpZ_{Pss}) which traverse the Hrp secretion pathway and that at least some PR-proteins can be induced by bacterial factors other than Hrp-controlled extracellular proteins.

Results

Symptoms on cucumber leaves treated with SAR inducers

Treatment of leaves with spores of C. lagenarium (a virulent, necrogenic pathogen of cucumber) resulted in the development of symptoms typically obtained with the fungus in cucumber, infiltrated areas were asymptomatic for 3-4 days, after which time tissues began to collapse and become necrotic. Lesions continued to expand for several days and developed a tan to brown pigmentation. Symptoms induced by treatments with Pss61 (an avirulent, HR necrosis-inducing pathogen) and HrpZ_{Pss} varied with environmental conditions in the greenhouse. Under high levels of natural light, Pss61 and HrpZ_{Pss} triggered the HR within 24 and 48 h, respectively, after infiltration. The HR was restricted to infiltrated areas and did not expand as did the necroses caused by C. lagenarium. Under lower natural light levels (cloudy days), tissues infiltrated with Pss61 or HrpZ_{Pss} developed a weaker HR characterized by increasing chlorosis over a 3-5 day period, then necroses developed gradually and irregularly, despite supplemental illumination with sodium lamps. Infiltration with hrpH (which is defective in the secretion of HrpZps. He et al., 1993) caused either no symptoms or a very mild chlorosis under all conditions tested. Infiltration with buffer alone caused only a small ring of white necrosis resulting from mechanical damage caused by pressure of the pipette mouth against the leaf. Interestingly, Infiltration with E amylovora harpin protein, which was prepared from DH5α(pCPP50) (He et al., 1994) and which induced a strong HR in tobacco leaves, did not induce HR necrosis in cucumber leaves (data not shown).

SAR to C. lagenarium

We first tested to see whether HrpZ_{Pss} alone could induce SAR to a well-studied fungal pathogen of cucumber, C. lagenarium. As shown in Table 1, HrpZ_{Pss} treatment induced SAR comparable to that induced by C. lagenarium

(approximately 90% reduction in total necrotic area relative to buffer-treated controls) in two upper leaves which expanded subsequent to induction treatment. The degrees of SAR induced by HrpZ_{Pss}, Pss61, Pss61-hrpH and C. lagenarium in cucumber were subsequently compared. Under conditions conducive to HR development in the greenhouse (high levels of natural light due to sunny weather) both HrpZ_{Pss} and Pss61 efficiently induced SAR in Leaf 2 and Leaf 3 (Table 2 and Figure 1a and b). SAR was expressed as a reduction in both the number and diameter of necrotic lesions resulting from challenge with C. lagenarium. Protection of Leaf 2 was comparable to that induced by C. lagenarium, whereas protection in Leaf 3 was weaker than that induced by the fungus. Under the conditions of this experiment, expansion of Leaf 2 and Leaf 3 occurred after the onset of the HR and necrosis incited by C. lagenarium Infiltration. Leaf 2 was fully expanded prior to challenge-inoculation, whereas Leaf 3 was not. The hrpH mutant did not induce SAR (Table 2). The quality and/or quantity of light profoundly influenced the induction of both the HR and SAR in cucumber by Pas61 and HrpZ_{Pss} in the greenhouse. When a similar experiment was conducted under conditions non-conducive to HR development (low levels of natural light on cloudy days), neither Pss61 nor HrpZps induced the HR or SAR, although C. lagenarium Incited necrotic lesions on Leaf 1 and induced SAR under these conditions (data not shown).

SAR to TNV

We next examined whether HrpZpss-induced SAR would be effective against a viral pathogen. In two initial experiments, the abilities of HrpZ_{Pas} and C. lagenarium to induce SAR to TNV were compared. HrpZ_{Psx} elicited a normal HR in these experiments and induced-SAR to TNV local lesion formation comparable to that induced by C. lagenarium (Table 3 and Figure 1c and d). We then compared the abilities of HrpZ_{Pss}, Pss81, hrpH, and C. lagenarium to induce SAR to TNV. Under high light conditions, HrpZ_{Pss} and Pss61 elicited a normal HR and induced SAR which restricted local lesion formation by TNV to an extent similar to that of SAR induced by C. lagenarium. The percentage of lesion number reduction was 68% for Pss61, 67.1% for HrpZ_{Pas}, and 75.5% for C. lagenarium (Table 3). Under low natural light conditions unfavorable for HR development (see Experimental procedures), HrpZ_{Pss} and Pss61 elicited a weaker degree of SAR relative to that induced by C. lagenarium. The percentage of lesion number reduction was 44.9% for Pss61, 46.7% for HrpZpss, and 89.6% for C. lagenarium (Table 3). The lesion numbers observed in these independent experiments varied greatly, mainly due to the use of different TNV inoculum preparations. TNV inoculum was prepared freshly each time from cucumber

Table 1. Induction by HrpZ_{Pss} and the fungal pathogen, C. lagenarium, of systemic acquired resistance to C. lagenarium in cucumber

		Lesf 2			Leaf 3		
Y	Lesion	Lesion diameter (mm)	Total necrotic area (mm²)	Lesion number	Lesion diameter (mm)	Total necrotic area (mm²)	
reatment uffer ImZ _{PM}	18.8 ± 0.5° 6.5 ± 0.9 3.3 ± 0.8	2.0 ± 0.1 1.1 ± 0.0 1.0 ± 0.0	60.9 ± 7.4 6.9 ± 1.3 2.6 ± 0.8	18.5 ± 0.6 9.5 ± 1.7 6.5 ± 1.3	2.5 ± 0.3 1.3 ± 0.1 1.2 ± 0.1	110.2 ± 29.0 13.4 ± 3.7 7.5 ± 1.3	

Leaf 1 of young plants was infiltrated with buffer (5 mM MgSO₄), or HrpZ_{Pas} (80 µg ml⁻¹) in buffer, or spores of C. lagenarium (5×10⁴ spores ml-1). After 7 days, Leaf 2 and Leaf 3 were challenged with 20 droplets per leaf containing spores of C. lagenarium. Disease was allowed to develop for 8 days.

Table 2 induction of systemic acquired resistance to C. lagenarium in cucumber by P. s. pv. syringse 61 (Pas61), HrpZ_{***} the hrpH mutant of Pas61 and C. lagenarium

		Leaf 2	. •		Leaf 3	
	Lesion number	Lesion diameter (mm)	Total necrotic area (mm²)	Lesion number	Lesion diameter - (mm)	Total necrotic area (mm²
Freatment Buffer hrpH Pse61 HrpZ-m C. legenarium	15.4±1.2° 13.2±1.1 6.4±0.4 5.0±0.5 4.0±1.2	1.6±0.2 1.7±0.1 1.2±0.1 1.2±0.1	38.9±8.3 32.1±2.2 7.0±1.9 5.9±1.4 8.4±5.3	16.2±1.0 15.4±1.6 9.4±1.1 8.6±2.5 6.4±1.4	1.8±0.1 1.8±0.1 1.5±0.1 1.6±0.2 1.4±0.2	52.0±8.1 50.0±11.9 21.2±6.2 24.4±9.1 13.2±5.0

Leaf 1 of young plants was infiltrated with buffer (5 mM MgSO₄), bacteria (OD₆₀₀=0.2), HrpZ_{Pas} (160 μg ml⁻¹), or spores of C. legenarium (5x10⁴ ml⁻¹). After 8 days, Leaf 2 and Leaf 3 were challenged with 20 droplets per leaf containing spores of *C. lagenarium*. Disease was allowed to develop for 8 days.

leaves bearing TNV lesions. In experiment 3, the hrpH mutant induced a low level of SAR to TNV (Table 3).

SAR to P. syringae pv. lacrymans

HrpZ_{Pss} and C. lagenarium also induced SAR to the angular leaf spot bacterium, P. s. pv. lecrymans. For these experiments, cucumber plants were challenge-inoculated at 11 days (by spraying) or 17 days (by rubbing) after treatment of Leaf 1 (Table 4). Although C. lagenarium was a more effective treatment, HrpZ_{Pas} also induced significant levels of SAR to the bacterium, reducing necrotic lesion numbers by 32 and 75%, compared with 50 and 86% for C. lagenarium, in the two experiments, respectively.

Induction of PR proteins

PR proteins that accumulated in treated cucumber plants were first analyzed using native polyacrylamide gel electrophoresis (PAGE). All treatments (C. lagenarium, Pss61 and HrpZ_{Pss} that induced SAR also induced the accumulation of three PR protein bands (tentatively named PR-A, PR-8 and PR-C) (Figure 2a). C. lagenarium induced PR-C, but not PR-A and PR-B, in systemic leaves, while Pss61 and HrpZ_{Pss} induced PR-B, but not PR-A and PR-C, in systemic leaves. Treatment with buffer or hrpH mutant did not induce these particular PR protein bands to levels that would allow visual identification. To see whether any PR proteins with known functions were induced in these plants, protein extracts were analyzed using native PAGE coupled with enzyme (chitinase, peroxidase and β -glucanase) activity staining. As shown in Figure 2(b), all three enzymes were induced in plants treated with HrpZPsp. Pss61 or C. lagenarium in both local (treated) and systemic leaves, although induction of chimnase isoforms by Pas61 and HrpZ_{Pas} in systemic leaves was variable and low. The enzyme activities were substantially higher in local leaves than in systemic leaves. Surprisingly, although the hrpH mutant bacterium failed to induce SAR, it efficiently induced peroxidase and chitinase, especially in treated leaves (Figure 2b). Only β-glucanese was not found to be induced to high levels in the hrpH-treated plants (Figure 2b). It is interesting to note that PR protein levels induced by various treatments correlated well with degrees of SAR induced by the same treatments (C. lagenarium >HrpZ_{Pss}=Pss61>hrpH> or = buffer).

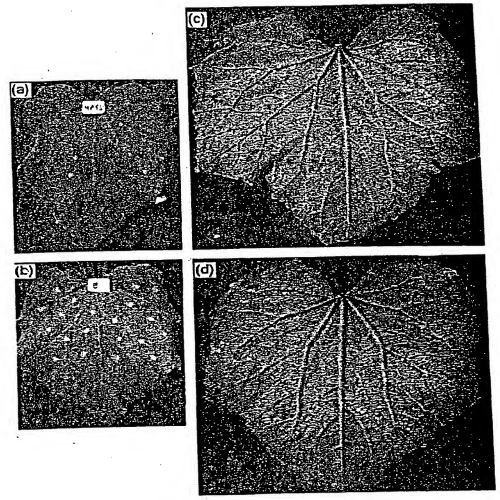


Figure 1. Disease symptoms caused by challenge-infection of *C. legensrium* and tobacco necrosis virus on cucumber leaves with or without prior induction of SAR.

Anthracnosa symptoms on Lasf 2 of cucumber plants with Lasf 1 previously treated with HrpZ_{pm} (80 µg mi⁻¹, a) or buffer (5 mM MgSO₄, b). Lasf 1 of young Anthracnosa symptoms on Lasf 2 of cucumber plants was infiltrated with buffer or HrpZ_{pm}. After 8 days, Lasf 2 and Lasf 3 were challenged with 20 droplets per lasf containing spores of *C. legenarium*. Disease was allowed to develop for 8 days, when the picture was taken.

TNV symptoms on Leaf 3 of cucumber plants with Leaf 1 previously treated with HrpZ_{Pax} (c) or buffer (d). Leaf 1 was treated by infiltration of buffer or HrpZ_{Pax} as described in footnotes to Table 1, After 7 days, Leaf 3 was challenged by mechanical inoculation with a TNV suspension prepared from infected cucumber leaves. Disease was allowed to develop for 9 days, when the picture was taken.

Induction of the pr-1 gene and SAR in tobacco

HrpZ_{Pss} also induced SAR to tobacco mosaic virus (TMV) in tobacco (Table 5). The SAR level induced by HrpZ_{Pss} was less than that induced by TMV. This was consistent with the different levels of induction of the *pr*-1 gene by HrpZ_{Pss} and TMV (Figure 3). TMV-inoculated local leaves (the third and fourth true leaves) also showed more necrosis than those infiltrated with HrpZ_{Pss} (data not shown), which may

be partly responsible for the different levels of SAR and pr-1 expression in TMV- and $HrpZ_{Psr}$ -induced plants.

Discussion

In this study, we show that $HrpZ_{Pss}$, a bacterial hrp gane product secreted via the Hrp pathway of P. s. pv. syringae, induced SAR in cucumber and tobacco. In cucumber, the

Table 3. Induction of systemic acquired resistance to TNV in cucumber by hrpH mutant, HrpZ_{Pas}, Pss81 and C. legenarium

	Number of TNV necrotic local lesions					
	Experiment 1	Experiment 2	Experiment 3	Experiment 4		
reatment	99.7±1 9.6 °	47.2±0.9b	730.0±63.9° 556.0±53.4	342.8±34.3° 324.3±11.2		
pH '	- 28.7±3.8	7.5±1.2	240.4±27.5	182.8±18.8 189.0±41.9		
IrpZ _{Pss} *se61 C. <i>lagenarium</i>	- 34.7±16.6	- 8.0±1.8	239.9±59.7 178.8±25.9	35.8±4.6		

*Mean ± SE of three replicate plants per treatment. **Mean ± SE of eight replicate plants per treatment. Leaf 1 was treated by infiltration of candidate inducers as described in the footnotes of Table 1. After 7 days, leaf 3 was challenged by mechanical inoculation with a TNV suspension prepared from infected cucumber leaves. Disease was allowed to develop for 10 or 9 days in experiments 1 and 2, respectively.

Experiments 1, 2 and 3 were performed under high levels of natural light during induction periods. Experiment 4 was performed on cloudy days.

Table 4, induction of systemic acquired resistance to P. syringae pv. lecrymans by HrpZ_{Pm} and C. lagenarium

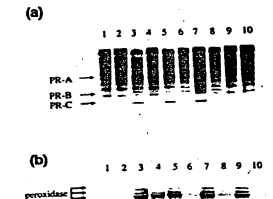
	Number of necrotic lesions*			
Treatment	Inoculated by rubbing	tnoculated by spraying		
Buffer	244.8±34.2	56.6±5.9		
HrpZ _{PM} C. lagenarium	168.5±24.5 122.8±9.8	13.8±1.7 8.3±2.1		

*Mean ± SE of five replicate plants per treatment. Leaf 1 of young plants was infiltrated with treatments as described in the footnotes of Table 1. Leaf 5 was challenged by rubbing, or by spraying the abaxial leaf surface with a suspension of bacterial cells (ODeco=0.2, 17 days after induction; or ODece=0.1, 11 days after induction, respectively). Disease was allowed to develop for 7 or 13 days in rub-inoculated or spray-inoculated plants, respectively.

efficacy against fungal, viral and bacterial pathogens and persistence (for at least 17 days in the bacterial challenge experiments) of HrpZ_{Pas}-induced SAR is comparable to that Induced by the bacterium (Pss61) that produces HrpZpss-The degree of SAR induced in cucumber by HrpZpm was also comparable to that induced by a well-studied biological inducer of SAR, C. lagenarium (Kuc and Richmond, 1977). The hrpH mutant of P. s. pv. syringse, which is defective in the secretion of HrpZPss and other proteinaceous pathogenicity factors (He et al., 1993; Huang et al., 1992; Yuan et al., in preparation), failed to induce SAR in cucumber. The induced PR protein patterns were the same in cucumber plants treated with Pss61 and HrpZPss, but were different from that in C. lagenarium-treated plants. Moreover, the hrpH mutant, although unable to induce SAR, efficiently induced at least two well-characterized PR proteins, chitinase and peroxidase (Figure 2b). These results suggest that the biological induction of SAR and PR proteins by P. s. pv. syringse 61 in the non-host plant, cucumber, is dependent on the production and secretion of proteinaceous elicitors of the HR, such as $\text{Hrp} Z_{\text{Pax}}$ but that at least some PR proteins can be induced by bacterial molecules independent of hrp gene functions.

The efficacy of both HrpZPs and Pss61 as inducers of SAR in cucumber appeared to be contingent upon their ability to elicit a normal HR, as low levels of natural light during the induction period, which interfered with HR development, resulted in reduced SAR to TNV and no SAR to C. legenarium (Table 3; Strobel and He, unpublished work). The negative effect of low light likely resulted from an effect on HR development rather than upon the plant's capacity to express SAR because C. lagenarium formed necrotic lesions typical of this compatible pathogen on Leaf 1 (the inducer leaf) and triggered SAR under these same conditions. The profound effect of light on the development of the HR has been observed previously (Sequeira, 1979), although the underlying mechanism remains to be determined. The dependence of the induction of SAR on the HR is further suggested by our observations that the hrpH mutant of Pss61, which produces but does not secrete HR elicitors (He et al.; 1993), did not elicit the HR or induce SAR in cucumber. Furthermore, E. amylovora harpin, another HR elicitor which is structurally different from $HrpZ_{Pss}$ and which elicited a strong HR in tobacco, did not induce an HR or SAR in cucumber plants (Strobel and He, unpublished observation). In conclusion, there appears to be a tight linkage between HR development and induction of SAR in plants by avirulent bacteria.

The tight linkage between the HR and SAR suggests that the signal(s) for the induction of SAR by HrpZ_{Pss} and P. s. pv. syringse 61 likely comes from dying plant cells and/or cells immediately adjacent to the dying cells during the HR. What types of cell death would lead to the induction of SAR? It has been shown that the HR triggered by live bacteria (Keen et al., 1981), HrpZ_{Pss} (He et al., 1993) or E. amylovora harpin (He et al., 1994) involves an active cell death pathway. Does this mean that only cells undergoing active cell death give rise to signals for SAR? The answer to this is probably not simple. SAR and PR proteins can



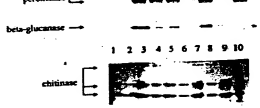


Figure 2. PR protein accumulation in cucumber plants.

PAGE (a) and PAGE coupled with activity staining (b) analyses of protein extracts from treated (lanes 1, 3, 5, 7 and 9) or systemic leaves (lanes 2, 4, 6, 8, and 10). The treatments were buffer (lanes 1 and 2), C. lagenarium (lanes 3 and 4), Pas51 (lanes 5 and 8), NrpZ_{Pa}, (lanes 7 and 8) and the hpH mutant (lanes 9 and 10), PR-A, PR-B and PR-C are tentative names for the three PR proteins observed in these experiments. The identities of these PR proteins are unknown.

be induced not only by HR-eliciting avirulent pathogens, but also by necrosis-causing virulent pathogens. For example, P. s. pv. lacrymans and C. lagenarium can efficiently induce SAR and/or PR proteins in the susceptible host plant, cucumber (Kuc and Richmond, 1977; Smith et al., 1991; this study). Unless cell death during the HR and some diseases shares the same biochemical processes, which is possible, the ability of both virulent and avirulent pathogens to induce SAR argues for multiple cell death pathways in the induction of SAR. On the other hand, not all types of plant cell death induce SAR. For example, cell death due to mechanical wounding or resulting from certain plant mutations does not induce SAR (Dietrich et al., 1994), it would be important in the future to learn why certain cell death processes, but not others, lead to SAR. Endogenous signaling molecules, such as salicylic acid and H₂O₂, have been shown or suggested to be involved in the induction of SAR (Chen et al., 1993; Gatiney et al., 1993; Malamy et al., 1990; Metraux et al., 1990; Rasmussen et al., 1991). However, the mechanism(s) by which various biological inducers of SAR generate these signals and the identity of the actual systemic signal(s) translocated from the induced leaves to distant leaves remain to be deter-

Table 5. Induction of systemic acquired resistance to TMIV by $HrpZ_{Pss}$ and TMIV

-	Diameter of necrotic lesions [®]		
Buffer	4.41±0.05		
HrpZ _{Pss}	3.05±0.03		
TMV	(2.34±0.03		

*Mean ± SE of 100 lesions per treatment.

The third and fourth true leaves of 6-week-old tobacco plants were inoculated with TMV (100-150 lesions per leaf), or infiltrated with 120 µg ml⁻¹ harpin_{Pss} or 5 mM MgSO₄ at 10 sites (50 µl per site). Five days later the seventh and eighth true leaves were challenge-inoculated with TMV. The diameters of TMV lesions on the challenged leaves were recorded.

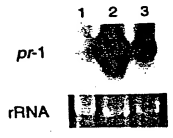


Figure 3. Induction of the pr-1 gene in tobacco leaves.

Total RNA was isolated from systemic leaves (the ninth true leaves) of plants treated with buffer (lane 1). TMV (lane 2), or HrpZ_{Pas} (lane 3) 5 days post-induction, A PCR-emplified internal fragment of the tobacco pr-1 gene was labeled with (c-³²P)IdATP and used as a probe. The largest rRNA species visualized after staining with enhichten bromide was used as a reference.

mined. Also, it has not been unequivocally shown that cell death is necessary for the induction of SAR.

It is interesting to observe that, sithough C. lagenarium (a necrotizing pathogen of cucumber), Pss61 (an HR-eliching bacterium on cucumber) and HrpZ_{Pas} (an HR-eliciting protein) all induced SAR in cucumber plants, there were some differences in the induction of PR proteins by these pathogens/protein. While C. lagenarium, Pss61 and HrpZpss all induced PR-A, PR-B and PR-C in the inoculated leaves, only C. lagenarium induced PR-C in systemic leaves to a high level (visible on a PAGE gel). In contrast, PR-B was induced in systemic leaves to high levels only by HrpZpas and Pss61. The induction patterns of PR-A, PR-B, PR-C, chitinase, peroxidase and β-glucanase were the same for Pss61 and HrpZ_{Pss}, suggesting that HrpZ_{Pss} either is a major inducer of SAR in Pss61 or is representative of SAR inducers produced by Pss61. The differences in the induction of PR proteins by C. lagenarium and Pse61/ HrpZ_{Pss} may have resulted from different inducers produced by C. lagenarium and Pss61/HrpZPss respectively. Alternatively, the differences may reflect possible mechanistic differences of plant cell death resulting from the HR caused by Pss61 or HrpZ_{Pss} and disease necrosis caused by C. lagenarium, respectively, although both types of cell death efficiently trigger SAR in cucumber.

In this study, 80-160 µg ml-1 purified HrpZ_{Pas} were used for induction of SAR. HrpZPss at these concentrations consistently elicited both HR and SAR in cucumber and tobacco leaves. It is not known whether these concentrations are comparable to the in vivo amounts of HrpZpm secreted by Pss61. Nor is it known whether the relative activity of purified HrpZ_{Pss} is comparable to that of HrpZ_{Pss} produced by Pss61 in planta. Previously, it was shown that Pss61 hrpZ mutants carrying transposon-induced mutations in the hrpZ gene (complementation group XII) were defective in the elicitation of HR (Huang et al., 1991) and SAR (data not shown). More recently, it was discovered that these transposon-induced hrpZ mutations exert a polar effect on five downstream hrp genes (hrpB-F) in the hrpZ operon (Preston et al., 1995; Collmer, personal communication). hrpB-F, like hrpH, are likely involved in the assembly of the Hrp secretion apparatus (Preston et al., 1995). Therefore, current hrpZ mutations affect the expression of not only the hrpZ gene but also several other hrp genes that are involved in the secretion of HrpZ_{Pss} and, most likely, other HR elicitors/pathogenicity factors. A non-polar hrpZ mutant is needed to assess the contribution of HrpZ_{Pss} in the induction of HR and SAR. Recently, several additional proteins traversing the P. syringae Hrp secretion pathway have been identified in P. syringae pv. tomato (Yuan et al., in preparation). It would be interesting to know whether some of these new Hrp-controlled P. syringae extracellular proteins can elicit HR and/or SAR.

Although the hrpH mutant of Pss61 failed to induce SAR in most experiments, it efficiently induced the accumulation of peroxidase and chitinase in all experiments (Figure 2b and data not shown). The induction of chitinasa by hrp mutants was also observed by Jakobek and Lindgren (1993). These data suggest that induction of PR proteins is not necessarily a reflection of induction of SAR and that the accumulation of certain PR proteins may not contribute to resistence. In our experiments, only the accumulation of β -glucanase seemed to correlate with the SAR induced by both C. lagenarium and Pss61/HrpZPss in cucumber. None of the other identified PR proteins were present at high levels in systemic leaves of all cucumber plants that exhibited SAR. Whether β-glucanese is responsible for the resistance of the induced plants to C. lagenarium, TNV and P. s. pv. lacrymans in cucumber remains to be investigated. The relationships between the PR-A, PR-B, and PR-C proteins with β -glucanese, chitinase, or peroxidase are not known.

The demonstration of HrpZ_{Pss} as a proteinaceous inducer of SAR may have important practical implications for plant disease management. Crop plants could be genetically

engineered with genes encoding proteinaceous HR/SAR inducers, such as HrpZ_{Pss} uncler the control of plant promoters inducible by virulent pathogens. If this approach were successful, the HR and SAR would be triggered in otherwise compatible interactions, limiting the disease development.

Experimental procedures

Growth of plants

Cucumber (Cucumis sativus L.) planets were grown in plastic pots containing Promix soil. A liquid fortilizer (Peter's 15-16-17, W. R. Grace and Co., Fogelsville, PA), containing 110 p.p.m. nitrogen, was supplied to the water, beginning when the first true leaf was fully open. Plants were grown in as glass greenhouse equipped with high-pressure sodium lights (with a photoperiod of 14 h) to supplement sunlight when necessary.

Preparation of inocula

HrpZha was purified by affinity chromatography from Escherichia coli DH5a(pSYH45). pSYH45 is a derivative of pOE30 (Qiagen, Inc.) expressing a hexahistidine-HrpZ_{Pm} (full-length) fusion protein. The first methionine residue of HrpZps was replaced by the following amino acid sequence in the fusion protein: MRGSHHHHHH. The fusion proteins was purified according to the manufacturer's instructions. Imidazole (300 mm) was used to elute HrpZ_{Pss} protein, followed by extensive dialysis (3000-fold) in 6 mM MgCl₂ at 4°C. The purity of HrpZ_{Pan} fusion protein was estimated by SDS-PAGE analysis to be greater than 95%. The fusion protein at the concentration of 80 µg ml⁻¹ elicited a strong HR in tobacco and cucumber leaves, while an identical preparation from DHSc(pQE30) (used as a control in the purification) did not elicit any visible response in the same leaves

Pseudomonas syringae strains were grown in King's B broth (King et al., 1954) overnight at 30°C. Bacterial suspensions were prepared in 5 mM MgSO. Spores of Colletotrichum lagenarium were prepared as described previously (Kuc and Richmond, 1977). Tobacco necrosis virus inoculum was prepared by grinding cucumber leaves bearing necrotic local leaions in water (1g infected leaf tissue per 10 ml distilled water).

Induction of SAR

First true leaves (Leaf 1) of young cucumber plants (cv. 'Marketer') were treated with test agents by infiltration through their abadial surfaces at 30 sites per leaf, with 10 µl per site delivered by a repeating pipettor. Treatments consisted of buffer (5 mM MgSQ₄), HrpZ_{Pss} (final concentration in buffer was 80-160 µg mf⁻¹), Pss61 or hrpH (a final ODecc=0.2 in 5 mM MgSO., equivalent to approximately 2×108 cells mil-1), or a spore suspension of C. lagenarium (7.5×10⁴ spores ml⁻¹).

For experiments involving tobacco (Nicotiana tabacum Samsun NN) plants, the third and fourth true leaves of 6-week-old plants were inoculated with TMV (100-150 lesions per leaf) or infiltrated with 120 µg mi⁻¹ HrpZ_{Pss} or 5 mM MgSO₄. For TMV inoculation, adaxial leaf surfaces were dusted with carborundum and then rubbed with a cheaseclotti pad moistened with a TMV suspension. For inoculation with HrpZ_{Pss} or 5 mM MgSO₄, 50 µl solution was pressured into each of 10 panels of a tobacco leaf using a needleless syringe. Five plants were used for each treatment.

Assessment of SAR

At 7-8 days after treatment of Leaf 1 with test agents, subsequently developed leaves (usually Leaf 2 and/or Leaf 3) were challenged with C. lagenarium, TNV or P. s. pv. lacrymans.

For fungal challenge, 20 sites per leaf received 10 µl droplets of a C. lagenarium spore suspension (1×10⁵ spores mir⁻¹) placed on adaxial surfaces with a repeating pipettor. After inoculation, plants were held in darkened moist chambers for 24 h to facilitate penetration of leaves by the pathogen. Chambers were then gradually opened to allow plant adaptation to ambient conditions over a 12 h period, and plants were then returned to a greenhouse bench for an additional 6-7 days to allow disease development.

For TNV challenge, adaxial leaf surfaces were dusted with carbonundum and then rubbed with a cheesecloth ped moistened with a TNV suspension. Virus-inoculated plants were maintained on a greenhouse bench for 8–10 days to permit disease development.

For assessment of SAR to the angular leaf spot bacterium, P. s. pv. lacrymans, Leaf 1 was infiltrated with buffer, C. lagenarium, or HrpZ_{Pse} as described above, and Leaf 5 was challenged on the abaxial surface with the becterium by spraying with a bacterial suspension (OD₆₀₀=0.1) containing 0.02% Silwet L-77, a surfactant, at 11 days post-induction or by rubbing with a cheesecloth pad asturated with a bacterial suspension (OD₆₀₀=0.2) at 17 days after induction treatment. Spray-inoculated leaves were misted once and plants were then placed in a darkened moist chamber for 18 h, followed by a 12 h acclimation period. Plants were subsequently returned to the greenhouse bench. Rub-inoculated leaves were misted once with water and plants were kept on a greenhouse bench. Disease was allowed to develop for 7 days for rub-inoculated plants or 13 days for spray-inoculated plants.

For evaluation of anthracnose development, the number and diameter of necrotic lesions caused by C. lagenarium were determined, and the total necrotic area per leaf was calculated. The extent of disease caused by TNV or P. s. pv. lacrymane was evaluated by counting necrotic local lesions on entire inoculated

For assessment of SAR to TMV, the seventh and eighth true leaves were challenge-inoculated with TMV (100–150 lesions per leaf) 5 days after induction. For each treatment the diameters of 100 TMV lesions (from 10 leaves of five plants) were recorded.

PR protein assay

Tissues were collected from Leaf 1 and Leaf 2 during the 14 day period following induction of Leaf 1. The leaf tissues were rapidly frozen with dry ice and stored at -80°C. Protein extraction was based on the method previously described (Ji and Kuc, 1995). Frozen leaf tissues were homogenized at 0-4°C in 0.1 M sodium citrate buffer, pH 5.A. containing 0.1% (w/v) β-mercaptoethanol and 0.1% (w/v) L-ascorbic acid. The homogenate was centrifuged at 12 000 g for 30 min. The supernatant was decanted and dialyzed against two changes of water for 24 h and then against two changes of 0.05 M sodium acetate buffer (pH 5.0) for 2 h. The extract was centrifuged again at 10 000 g for 10 min. The supernatant was used as crude enzyme extract. Protein concentrations were measured using the Bio-Rad protein assay kit with bovine gamma globulin as standard.

Determination of enzyme activities in cucumber leaves

Protein patterns and peroxidase isozymes were analyzed after a single separation using a 15% (w/v) native-PAGE gel (Pan et al., 1989). Peroxidase activity was determined using guaiscol as substrate (Hammerschmidt et al., 1982). β-1,3-glucanase and chitinase activities were detected as described elsewhere (Ji and Kuc, 1995).

Expression of pr-1 gene in tobacco leaves

An internal fragment (from nt 304 to 535) of the tobacco pr-1 gene (Figure 1 in Cornelissen et al., 1986) was amplified in a polymerase chain reaction (PCR) and labeled with (c. 32P)-dATP. Total RNA was purified from systemic leaves (the ninth true leaves) of tobacco plants 5 days post-induction. Ten micrograms of RNA from each treatment were fractionated in a 1.2% agarose/formaldehyde gel and subsequently blotted to Immobilion-N membrane (Millipore). Hybridization was performed in a solution consisting of 6×SSC, 2×Denhardt's reagent, 0.1% SDS and 10% dextran sulfate at 55°C. Washes were carried out in 0.2×SSC, 0.1% SDS at 60°C.

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(54) DISEASE-RESISTANT PLANTS AND METHOD OF CONSTRUCTING THE SAME

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(57) ABSTRACT

It is the object of the present invention to provide diseaseresistant plants which have been transformed to cause an effective defense reaction, and methods for producing the same.

The present invention provides expression cassettes comprising a promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression, and a gene, under the control of said promoter, encoding an elicitor protein.

Plants which the construct was introduced	Tobacco	Торассо	Rice, Tobacco	Rice, Tobacco	
Contents of the construct	PAL1.45 pro hrpZ	PAL0.45 pro hrpZ	35S pro hrpZ	PPDK pro hrpZ	
Inducible/ Constitutive	Inducible	Inducible	Constitutive	Constitutive	
Construct	PALL-hrpZ	PALS-hrpZ	35S-hrpZ	PPDK-hrpZ	

Fig. 1 Constructs introduced into plants

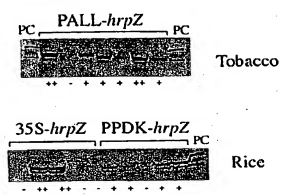
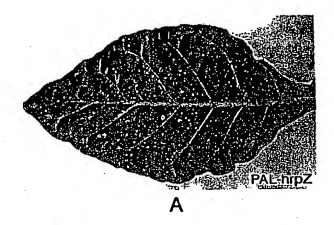


Fig. 2 Expression of harpin_{pss} in tobacco and rice



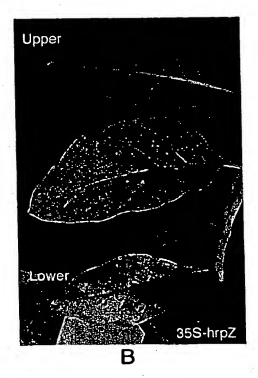


Fig. 3 Formation of hypersensitiveresponse-like localized necrosis spots

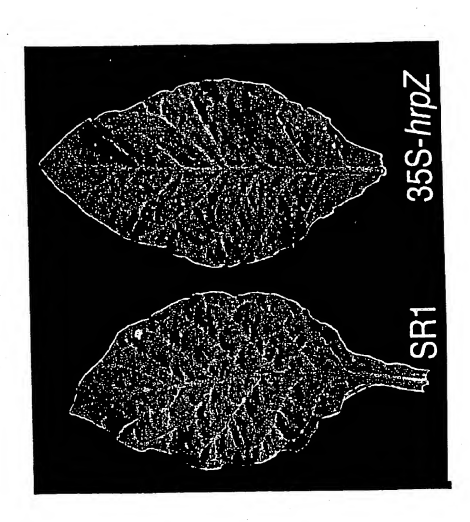


Fig. 4 Resistance to powdery mildew

DISEASE-RESISTANT PLANTS AND METHOD OF CONSTRUCTING THE SAME

FIELD OF THE INVENTION

[0001] The present invention relates to methods for producing disease-resistant plants, gene expression cassettes for producing disease-resistant plants, and transgenic, disease-resistant plants produced by the method.

BACKGROUND OF THE INVENTION

[0002] Plant defense against pathogens differs in its mechanism from that observed in animals. For example, there is known in higher plants a hypersensitive response (HR) mechanism which involves a dynamic resistance reaction to pathogen invasion. When a pathogen invades a plant, plant cells at a site of invasion die in response, whereby pathogens are trapped locally. This reaction is known to be induced as a result of either an incompatible host-pathogen interaction or a non-host-pathogen interaction. Such cell suicide can be understood in terms of a localized, programmed cell death (Dangl et al.: Plant Cell 8: 1973-1807 (1996)). In addition to the mechanism involving HR, other defense reactions, including generation of active oxygen species, reinforcement of a cell wall, production of phytoalexin and biosynthesis of defense-related proteins such as PR proteins, are also known (Hammond-Kosack and Jones: Plant Cell 8: 1773-1791 (1996)). Further, in addition to such localized defense responses, there is known to take place in many cases a defense reaction spreads whereby PR proteins accumulate also in non-infected parts of a plant, whereby resistance is imparted to the entire plant. This mechanism is referred to as systemic acquired resistance (SAR) and continues for several weeks or longer. As a result, the entire plant is made resistant to secondary infection (Sticher et al.: Annu. Rev. Phytopathol. 35: 235-270 (1997)).

[0003] A first reaction of a plant of switching on a highly organized defense reaction such as outlined above is the recognition by the plant of a molecule called an "elicitor" directly or indirectly produced by an invading pathogen. Additionally, complex signal cascades including the subsequent rapid generation of active oxygen species and reversible protein phosphorylation are considered to be important as initial reactions of the defense response (Yang et al.: Genes Dev. 11: 1621-1639 (1997)). There are a wide variety of elicitors, including so-called nonspecific elicitors e.g. oligosaccharides which are products by degradation of cell wall components of many fungi including chitin/chitosan and glucan, or oligogalacturonic acids derived from a plant cell wall, variety-specific elicitors e.g. avirulence gene products of pathogens such as AVR 9 (Avr gene products), and elicitors with an intermediate specificity such as elicitin (Boller: Annu. Rev. Plant Physiol. Plant Mol. Biol. 46: 189-214 (1995)).

[0004] Harpin is a bacterium-derived protein elicitor which induces hypersensitive cell death in a non-host plant (Wei et al.: Science 257: 85-88 (1992), He et al.: Cell 73: 1255-1266 (1993)). Harpin (harpin_{Ea}) has been purified as a first bacterium-derived HR-inducing protein from Erwinia amylovora Ea321, a pathogen of pear and apple, and Escherichia coli transformed with a cosmid containing the hrp gene cluster, and an hrpN gene encoding Harpin has been cloned (Wei et al.: Science 257: 85-88 (1992)). There-

after, harpingpss encoded by hrpZ gene has been identified and characterized from Pseudomonas syringae pv. syringae 61, a pathogen of a bean, by screening an Escherichia coli expression library with an activity of inducing HR to a tobacco leaf as an index (He et al.: Cell 73: 1255-1266 (1993), and Japanese Patent Application Domestic Announcement No. 1996-510127). The homology between these two harpins is low, and a relatively high homology is found only in 22 amino acids. Moreover, the role of a harpin in pathogenicity has not been made clear. In addition to these, as a third protein, PopA protein (which PopA encodes) is identified from Pseudomonas solanacearum GMI1000, a pathogen of a tomato, as a protein inducing HR to a non-host tobacco (Arlat et al.: EMBO. J. 13: 543-553 (1994)). Though PopA gene is located on the outside of hrp cluster, differing from hrpN and hrpZ, they are identical in that they are under the control of an hrp regulon. The above three proteins are glycine-rich, heat stable proteins, induce HR to a non-host tobacco and are secreted extracellularly at least in vitro in a manner of depending upon hrp protein. In addition to these are reported HrpW protein from Pseudomonas syringae pv. tomato DC3000 as a protein having the same function (Charkowski et al.: J. Bacteriol. 180: 5211-5217 (1998)), hrpZ_{ps} and hrpZ_{psg} proteins as harpin_{pss} homologues (Preston et al.: Mol. Plant-Microbe. Interact. 8: 717-732 (1995)), and harpin_{Ech} (Bauer et al.: Mol. Plant-Microbe. Interact. 8: 484-491 (1995)) and hrpN_{Ecc} protein (Cui et al.: Mol. Plant-Microbe. Interact. 9: 565-573 (1996)) as harping homologues.

[0005] It has been made apparent from studies upon various metabolic inhibitors that the formation of localized necrosis spots with harpin is not so-called necrosis due to the cytotoxicity of harpin but a cell death resulting from a positive response on the plant side (He et al.: Mol. Plant-Microbe. Interact. 7: 289-292 (1994), and He et al.: Cell 73: 1255-1266 (1993)), and this hypersensitive cell death is thought to be a type of programmed cell death (Desikan et al.: Biochem. J. 330: 115-120 (1998)). The addition of harpin into a cell culture of Arabidopsis induces a homologue of gp91-phox, a constituent of NADPH oxidase, which is thought to have an important role in the oxidative burst as an initial reaction of a disease-resistant reaction, (J. Exp. Bot. 49: 1767-1771 (1998)), and mitogen-activated protein (MAP) kinase (Desikan et al.: Planta. 210: 97-103 (1999)). Moreover, a harpin can impart systemic acquired resistance (SAR) to a plant. For example, SAR meditated by salicylic acid and an NIM gene can be induced to an Arabidopsis plant by artificially injecting harpinge into the plant cells (Dong et al.: The Plant J. 20: 207-215 (1999)), and Harpin can induce SAR to a cucumber and impart a wide spectrum of resistance to fungi, viruses and bacteria (Strobel et al.: Plant J. 9: 431-439 (1996)).

[0006] Thus, there are reports about artificially injecting or spraying purified harpin into a plant and analyzing the induction of a hypersensitive cell death and an acquired resistance reaction (Japanese Patent Application Domestic Announcement No. 1999-506938, Strobel et al.: Plant J. 9: 431-439 (1996), and Dong et al.: The Plant J. 20: 207-215 (1999)). However, there is no report about introducing a gene encoding an elicitor protein such as a harpin into a plant to produce a transgenic plant and analyzing it.

SUMMARY OF THE INVENTION

[0007] It has been anticipated that, when a gene encoding an elicitor protein such as harpin is introduced into a plant, the plant will express an elicitor protein at a certain amount, even in a normal state with no pathogen, or that it will also express an elicitor protein in a certain amount in organs other than those invaded with a disease, and as a result, various unintended reactions occur to prevent the plant from growing normally. The object of the present invention is therefore to provide a disease-resistant transgenic plant which has been transformed to induce a proper defense reaction, and to provide a method for producing the same.

[0008] The present inventors have engaged in studies assiduously, and as a result have found that a transgenic tobacco with hrpZ gene of Psedomonas syringae pv. syringae LOB2-1 introduced thereinto induces hypersensitiveresponse-like localized necrosis spots in response to the inoculation of a powdery mildew fungi (Erysiphe cichoracearum) to become resistant, which has led to the completion of the present invention. Surprisingly, a plant grew normally when cell-death-inducing harpin was expressed with a constitutive promoter (cauliflower mosaic virus 35S RNA gene promoter) capable of promoting expression in cells of the whole body. In addition, a hypersensitive celldeath-like reaction was induced only after inoculation with a pathogen. Further, the present inventors have found that a transgenic rice with the same hrpZ gene introduced thereinto becomes blast (Magnaporthe grisea)-resistant, thus showing the general-applicability of the present invention.

[0009] The present invention provides a transgenic, disease-resistant plant which has been transformed with an expression cassette comprising a promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression and a gene encoding an elicitor protein under the control of said promoter, wherein said plant is capable of effecting the constitutive, inducible, or organ- or phase-specific expression of the elicitor protein in an amount effective for inducing a defense reaction.

BRIEF DESCRIPTION OF THE DRAWINGS

[0010] FIG. 1 shows the constructs constructed and introduced into plants in the present invention.

[0011] FIG. 2 is a photograph showing exemplary of the detection results using Western analysis for harpin_{pas} accumulation in transgenic tobacco and rice of the T₀ generation. PC represents harpin_{pas} expression in *Escherichia coli* as a control.

[0012] FIG. 3 is a photograph showing the appearances of localized necrosis spots occurring in a transgenic tobacco of the T₁ generation. A: PALL-hrpZ-introduced individual (5th day after inoculation, harpin expression level: ++), B: 35S-hrpZ-introduced individual (7th day after inoculation, harpin expression level: ++)

[0013] FIG. 4 is a photograph showing the resistance of a transgenic tobacco of the T_1 generation against powdery mildew. (Right: 35S-hrpZ-introduced individual, harpin expression level: ++, Left: SR1 as a control, 11th day after inoculation in both)

DETAILED DESCRIPTION OF THE INVENTION

[0014] The present invention also provides methods for producing transgenic, disease-resistant plants capable of

effecting the constitutive, inducible, or organ- or phasespecific expression of an elicitor protein in an amount effective for inducing a defense reaction. Such methods comprise the steps of: (a) obtaining transgenic plant cells with expression cassettes comprising a promoter capable of promoting a constitutive, inducible, or organ- or phasespecific gene expression and a gene encoding an elicitor protein under the control of said promoter; and (b) regenerating a complete plant from said transgenic plant cell.

[0015] The present invention also provides expression cassettes capable of being employed for producing a transgenic, disease-resistant plants. Such expression cassettes comprise at least: (a) a promoter capable of promoting a constitutive, inducible, or organ or phase-specific gene expression; and (b) a gene, under the control of said promoter, encoding an elicitor protein. "Elicitor" is a general term used for substances inducing defense reactions in plants, and including heavy metal ions, and cell wall components of pathogens or plants, in addition to proteins. The term "elicitor" as used in the present specification refers to a protein elicitor unless otherwise specified.

[0016] The term "elicitor protein" as used in the present invention can be any protein capable of inducing a proper defense reaction in a plant to be transformed, and preferably a protein possessing a hypersensitive-response-inducing activity against pathogenic microorganisms. It includes harpin and a harpin-like protein having the same function as harpin. "Harpin" is a protein expected to be introduced into a plant in a manner of depending upon hrp gene though the Type III secretion mechanism, and includes, in addition to harpin_{pss}, (He et al.: Cell 73: 1255-1266 (1993), and Japanese Patent Application Domestic Announcement[kohyo] No. 510127/96), harpin_{Ea} (Wei et al.: Science 257: 85-88 (1992), and Japanese Patent Application Domestic Announcement[kohyo]No. 506938/99), PopA (Arlat et al.: EMBO. J. 13: 543-553 (1994)), and hrpW protein (Charkowski et al.: J. Bacteriol. 180: 5211-5217 (1998). Additionally the protein possessing a hypersensitive-response-inducing activity can be, for example, (a) a proteinconsisting of the amino acid sequence of SEQ. ID No. 2; (b) a protein consisting of an amino acid sequence derived from the amino acid sequence of SEQ. ID No. 2 by deletion, substitution, addition or insertion of one or more amino acids, and possessing a hypersensitive-response-inducing activity; or (c) a protein consisting of an amino acid sequence being at least 50% (preferably at least 80%, more preferably at least 90%, and still more preferably at least 97%) homologous to the amino acid sequence of SEQ. ID No. 2, and possessing a hypersensitive-response-inducing activity. A protein consisting of the amino acid of SEQ ID No. 2 is novel. Hence, the present invention provides one of the following proteins: (a) a protein consisting of the amino acid sequence of SEQ. ID No. 2; (b) a protein consisting of an amino acid sequence derived from the amino acid sequence of SEQ. ID No. 2 by deletion, substitution, addition or insertion of one or more amino acids, and possessing a hypersensitive-response-inducing activity; and (c) a protein consisting of an amino acid sequence being at least 97% homologous to the amino acid sequence of SEQ. ID No. 2, and possessing a hypersensitive-response-inducing activity (but known proteins themselves are excluded from the scope of the present invention).

[0017] By "Homology" referred to in connection with amino acid sequences in the present specification is meant a degree of identification of amino acid residues constituting each sequence between sequences to be compared. In homology, the existence of a gap(s) and the nature of an amino acid(s) are taken into consideration (Wilbur, Proc. Natl. Acad. Sci. USA 80: 726-730 (1983) and the like). To calculate homology, commercially available software such as BLAST (Altschul: J. Mol. Biol. 215: 403-410 (1990), and FASTA (Peasron: Methods in Enzymology 183: 63-69 (1990)) can be employed.

[0018] The description "deletion, substitution, addition or insertion of one or more amino acids" as used in the present specification in connection with an amino acid sequence in the means that a certain number of an amino acid(s) are substituted etc. by any well known technical method such as site-specific mutagenesis, or naturally. The number is, for example, up to ten, and is preferably from 3 to up to 5.

[0019] A gene encoding an elicitor protein to be employed in the expression cassette of the present invention can easily be isolated by methods well-known to those skilled in the art.

[0020] The gene encoding an elicitor protein can be, for example, (a) a DNA molecule consisting of the nucleotide sequence of SEQ. ID No. 1; (b) a DNA molecule consisting of a nucleotide sequence derived from the nucleotide sequence of SEQ. ID No. 1 by deletion, substitution, addition or insertion of one or more nucleotides, and encoding a protein possessing a hypersensitive-response-inducing activity; (c) a DNA molecule consisting of a nucleotide sequence being hybridizable with a DNA molecule consisting of the nucleotide sequence complementary to the nucleotide sequence of SEQ. ID No. 1 under stringent conditions, and encoding a protein possessing a hypersensitive-response-inducing activity; or (d) a DNA molecule consisting of a nucleotide sequence being at least 50% (preferably at least 80%, more preferably at least 90%, and still more preferably at least 97%) homologous to the nucleotide sequence of SEQ. ID No. 1, and encoding a protein possessing a hypersensitive-response-inducing activity. A DNA molecule consisting of the nucleotide sequence of SEQ ID No. 1 is novel. Hence, the present invention also provides a gene consisting of one of the following DNA molecules: (a) a DNA molecule consisting of the nucleotide sequence of SEQ. ID No. 1; (b) a DNA molecule consisting of a nucleotide sequence derived from the nucleotide sequence of SEQ. ID No. 1 by deletion, substitution, addition or insertion of one or more nucleotides, and encoding a protein possessing a hypersensitive-response-inducing activity; (c) a DNA molecule consisting of a nucleotide sequence being hybridizable with a DNA molecule consisting of the complementary nucleotide sequence to the nucleotide sequence of SEQ. ID No. 1 under stringent conditions, and encoding a protein possessing a hypersensitive-response-inducing activity; or (d) a DNA molecule consisting of a nucleotide sequence being at least 50% homologous to the nucleotide sequence of SEQ. ID No. 1, and encoding a protein possessing a hypersensitive-response-inducing activity (but known genes themselves such as hrpZ gene of Pseudomonas syringae pv. syringae 61 are excluded from the scope of the present invention). To calculate homology in connection with nucleotide sequences, commercially available software can be employed.

[0021] By "deletion, substitution, addition or insertion of one or more nucleotides" in connection with a nucleotide sequence in the present specification is meant that a certain number of a nucleotide(s) are substituted etc. by a well-known technical method such as a site-specific mutagenesis or naturally. The number is, for example, up to ten, preferably from 3 to up to 5. By "stringent conditions" referred to in the present specification is meant hybridization conditions wherein the temperature is at about 40° C. or above and that the salt concentration is of about 6xSSC (1xSSC=15 mM sodium citrate buffer; pH: 7.0; 0.15 M sodium chloride; 0.1% SDS), preferably at about 50° C. or above, more preferably at about 65° C. or above.

[0022] The promoter to be employed in the present invention can be any promoter capable of functioning as a promoter for a gene encoding an elicitor protein in a plant to be transformed. In the present invention, a promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression can be employed.

[0023] By "promoter promoting a constitutive gene expression (often referred to as a "constitutive promoter")" is meant a promoter whose organ specificity and/or phase specificity are (is) not high in connection with the transcription of the gene. Examples of the constitutive promoter include cauliflower mosaic virus 35S promoter, ubiquitin promoter (Cornejo et al.: Plant Mol. Biol. 23: 567-581 (1993)), actin promoter (McElroy et al.: Plant Cell 2: 163-171 (1990)), alpha tubulin promoter (Carpenter et al.: Plant Mol. Biol. 21: 937-942 (1993)) and Sc promoter (Schenk et al.: Plant Mol. Biol. 39: 1221-1230 (1999)). In a transgenic plant, the expression cassette promoting the constitutive expression of an elicitor protein includes, for example, a known promoter that is known as a constitutive promoter.

[0024] By "promoter promoting an inducible gene expression (often referred to as an "inducible promoter")" is meant a promoter which induces transcription by physical or chemical stimulation, such as light, disease, injury or contact with an elicitor. Examples of the inducible promoter include pea PAL promoter, Prp1 promoter (Japanese Patent Application No. 1998-500312), hsr203J promoter (Pontier et al.: Plant J. 5: 507-521 (1994)), EAS4 promoter (Yin et al.: Plant Physiol. 115: 437-451 (1997)), PR1b1 promoter (Tornero et al.: Mol. Plant Microbe. Interact. 10: 624-634 (1997)), tap1 promoter (Mohan et al.: Plant Mol. Biol. 22: 475-490 (1993)) and AoPR1 promoter (Warner et al.: Plant J. 3: 191-201 (1993)). In a transgenic plant, the expression cassette promoting an inducible elicitor protein expression includes, for example, a known promoter known as an inducible promoter.

[0025] By "promoter promoting an organ-specific gene expression (often referred to as an "organ-specific promoter")" is meant a promoter giving, to the transcription of the gene, a specificity to an organ, such as a leaf, a root, a stem, a flower, a stamen and a pistil. Examples of the organ-specific promoter include a promoter promoting a high gene expression in green tissues of a photosynthesis-related gene, such as PPDK (Matsuoka et al.: Proc. Natl. Acad. Sci. USA 90: 9586-9590 (1993)), PEPC (Yanagisawa and Izui: J. Biochem. 106: 982-987 (1989) and Matsuoka et al.: Plant J. 6: 311-319 (1994)) and Rubisco (Matsuoka et al.: Plant J. 6: 311-319 (1994)). In a transgenic plant, the

expression cassette promoting an organ-specific elicitor protein expression includes, for example, a known promoter that is known as an organ-specific promoter.

[0026] By "promoter promoting a phase-specific gene expression (often referred to as a "phase-specific promoter")" is meant a promoter giving, to the transcription of the gene, a phase specificity to a phase, such as a initial, middle and later growth phase. Examples of the phase-specific promoter include a promoter functioning specifically in aged leaves such as SAG12 promoter (Gan and Amashino: Science 270: 1986-1988 (1985)).

[0027] Vectors for sub-cloning each DNA fragment as a component of the expression cassette of the present invention can be simply prepared by connecting an intended gene into a vector for recombination (plasmid DNA) available in the art by any common technique. Specific examples of suitable vectors include plasmids derived from *Escherichia coli*, such as pBluescript, pUC18, pUC19 and pBR322, but are not limited only to these plasmids.

As a vector for introducing the expression cassette of the present invention into a plant to be transformed, a vector for transforming plants can be used. The vectors for plants are not particularly limited, so far as they are capable of expressing the concerned gene and producing the concerned protein in a plant cell, and examples thereof include pBI221, pBI121 (both being manufactured by Clontech) and vectors derived therefrom. In addition, for the transformation of a monocotyledonous plant in particular, there can be exemplified pIG121Hm, pTOK233 (both by Hiei et al.: Plant J. 6: 271-282 (1994)), pSB424 (Komari et al.: Plant J. 10: 165-174 (1996)), superbinary vector pSB21 and vectors derived therefrom. A recombination vector having the expression cassette of the present invention can be constructed by introducing a gene encoding an elicitor protein into any of these known vectors (if required, a promoter region being recombined) by a procedure known well to those skilled in the art. For example, a recombinant vector having an expression cassette comprising a constitutive promoter and hrpZ gene can be constructed by integrating hrpZ gene into superbinary vector pSB21. A recombinant vector having an expression cassette comprising an inducible promoter and hrpZ gene can be constructed by removing the existing promoter from the above recombinant vector and integrating an inducible promoter in place.

[0029] A plant-transforming vector preferably comprises at least a promoter, a translation initiator codon, a desired gene (a DNA sequence of the invention of the present application or a part thereof), a translation termination codon and a terminator. Moreover, it may comprise a DNA molecule encoding a signal peptide, an enhancer sequence, a non-translation region on the 5' side and the 3' side of the desired gene and a selection marker region as appropriate. Examples of marker genes include antibiotic-resistant genes such as tetracyclin, ampicillin, kanamycin or neomycin, hygromycin or spectinomycin; and genes such as luciferase, β-galactosidase, β-glucuronidase(GUS), green fluorescence protein (GFP), β-lactamase and chloramphenicol acetyl transferase (CAT).

[0030] As methods for introducing a gene into a plant can be mentioned a method employing an agrobacterium (Horsch et al.: Science 227: 129 (1985), Hiei et al.: Plant J. 6: 271282 (1994)), a leaf disc method (Horsch et al.: Science

227: 1229-1231 (1985), an electroporation method (Fromm et al.: Nature 319: 791 (1986)), a PEG method (Paszkowski et al.: EMBO. J. 3: 2717 (1984)), a micro-injection method (Crossway et al.: Mol. Gen. Genet. 202: 179 (1986)) and a minute substance collision method (McCabe et al.: Bio/ Technology 6: 923 (1988)), but any method for introducing a gene into a desired plant may be employed without any particular limitation. Of these methods for transfection, a method comprising transferring a vector into an agrobacterium by mating and then infecting a plant with the agrobacterium is preferred. Methods for infection is also wellknown to those skilled in the art. Examples include a method comprising damaging a plant tissue and infecting it with a bacterium; a method comprising infecting an embryo tissue (including an immature embryo) of a plant with the bacterium; a method comprising infecting with a callus; a method comprising co-culturing protoplasts and the bacterium; and a method comprising culturing a fragment of a leaf tissue together with the bacterium (leaf disc method).

[0031] Successfully transformed cells can be selected from other cells by employing an appropriate marker as an index or examining the expression of a desired trait. The transformed cell can further be differentiated employing a conventional technique to obtain a desired transgenic plant.

[0032] Analysis of the resultant transformant can be performed by employing various methods that are well-known to those skilled in the art. For example, oligonucleotide primers can be synthesized according to the DNA sequence of the introduced gene, and the chromosome DNA of the transgenic plant can be analyzed by PCR employing the primers. In addition, the analysis can be performed on the basis of the existence of mRNA corresponding to the introduced gene and the existence of the protein expression. Moreover, the analysis can be performed on the basis of the appearance of the plant (for example, in the case of transformation with a gene encoding a protein capable of inducing localized necrosis spots, the presence of localized necrosis spots, or the size, number and the like of the localized necrosis spots), disease resistance (for example, the existence of resistance or its degree upon contacting the plant with a pathogen) and the like.

[0033] In the transgenic plant of the present invention, a constitutive, inducible, or organ- or phase-specific expression of an elicitor protein in an amount effective for inducing a defense reaction can be achieved. The amount effective for inducing a defense reaction is such an amount that the expressed elicitor protein can induce at least a localized defense-related reaction (for example, induction of a hypersensitive cell death (localized necrosis)) to the plant. Preferably, the amount is such that the defense reaction extends to the whole body of the plant, and as a result, the whole plant becomes resistant (systemic acquired disease-resistant). Moreover, preferably, the amount is not so large that causes death of the localized tissue having the necrosis spots as a result of the localized necrosis spots becoming too large.

[0034] Moreover, in the transgenic plant of the present invention, an elicitor protein is preferably expressed in an amount which, while being effective for inducing a defense reaction in response to stimulation such as the invasion of a pathogen, does not, under normal conditions, remarkably prevent the growth of the plant due to the negligible or low expression, if any. For example, in the case of employing

harpin_{pas} as an elicitor protein, usually no harpin_{pas} is expressed, or is expressed only in an amount that does not allow localized necrosis spots to cause the death of the organ, and preferably it is expressed in an amount that induces a hypersensitive response at the time of the invasion of a pathogen. Further, it is preferably expressed in such an amount that, even if a pathogen invades to cause harpin_{pas} to accumulate, localized necrosis spots are hardly observable by the naked eye, but the whole body acquires a systemic disease-resistannce.

[0035] In order to induce such a proper defense reaction, for example, a promoter capable of promoting an inducible gene expression is employed. Hence, in one embodiment of the present invention, an inducible promoter and a harpin gene are combined.

[0036] In addition, a proper defense reaction can be accomplished not only in the case of employing an inducible promoter but also in the case of employing a constitutive promoter. Hence, in another embodiment of the present invention, a constitutive promoter and a harpin gene are used in combination. In this embodiment, as a mechanism of the occurrence of a proper defense reaction, it is considered that an elicitor protein, for example, harpin, so, is recognized at the outside of cell membranes or on the cell wall of plant cells, and hence, harpin, accumulating in cytoplasm is not recognized by plant cells until degradation of cells occurs due to invasion of fungus, and as a result, the hypersensitive response appears after the inoculation of the pathogen or it is deduced that there exists a further factor which is related to the inoculation of a pathogen in the mechanism of the occurrence of the elicitor activity of harpin, as.

[0037] The transgenic plants of the present invention include a transgenic, powdery mildew-resistant tobacco which has been transformed with an expression cassette comprising a constitutive or inducible promoter and a gene, under the control of said promoter, encoding an elicitor protein such as harpin_{pes}, or a transgenic, blast-resistant rice which has been transformed with an expression cassette comprising a constitutive promoter and a gene, under the control of the promoter, encoding an elicitor protein such as harpin_{pes}.

[0038] It is thought that the present invention can be applied to plants other than rice and tobacco described in the examples to be described later. Examples of such plants include, as crops, wheat, barley, rye, corn, sugar cane, sorghum, cotton, sunflower, peanut, tomato, potato, sweet potato, pea, soybean, azuki bean, lettuce, cabbage, cauliflower, broccoli, turnip, radish, spinach, onion, carrot, eggplant, pumpkin, cucumber, apple, pear, melon, strawberry and burdock; and, as ornamental plants, arabidopsis thaliana, petunia, chrysanthemum, carnation, saintpaulia and zinnia. The "transgenic plants" referred to in the present invention include not only transgenic plants (To generation) obtained by obtaining a transgenic plant cell according to the method of the present invention and regenerating, from said plant cell, a complete plant, but also later-generation (T1 generation and the like) plants obtained from said transgenic plants so far as the disease-resistant trait is contained. In addition, the "plants" referred to in the present invention include, unless otherwise speified, in addition to plants (individuals), seeds (including germinated seeds and immature seeds), organs or parts thereof (including a leaf, a root,

a stem, a flower, a stamen, a pistil and pieces thereof), a plant culture cell, a callus and a protoplast.

[0039] The diseases analyzed in the following examples are tobacco powdery mildew and rice blast, but as other diseases of tobacco there can be mentioned wildfire, bacterial wilt and TMV; and as other diseases of rice there can be mentioned sheath blight disease and bacterial leaf blight disease. According to the method for producing a disease-resistant plant of the present invention, it is possible to impart resistance in plants to these diseases.

EXAMPLES

Example 1

Cloning of HrpZ Gene

[0040] A pair of primers for amplifying the open leading frame of hrpZ gene were synthesized in reference to the nucleotide sequence of the reported hrpZ gene of *Pseudomonas syringae* pv. syringae 61 (He et al.: Cell 73: 1255-1266 (1993)), and Japanese Patent Application Domestic Announcement[Kohyo] No. 1996-510127):

Hrp1: AAA ATC TAG AAT GCA GAG TCT CAG TCT TAA

Hrp2: AAA AGT CGA CTC AGG CTG CAG CCT GAT TGC

[0041] Employing these primers, PCR was performed with a DNA molecule of a cosmid clone containing an hrp cluster derived from Pseudomonas syringae pv. syringae LOB2-1 (a casual agent for bacterial blight of lilac) (Inoue and Takikawa: J. Gen. Plant Pathol. 66: 238-241 (2000)) as a template. PCR was performed under the following conditions: the amount of a reaction solution: 20>1; each primer: 0.5 μM; dNTP: 0.2 mM; 1×ExTaq buffer; ExTaq DNA polymerase (from Takara Shuzo): 1U; once at 95° C. for 5 minutes, then 30 cycles at 94° C. for 30 seconds, at 60° C. for 30 seconds and at 72° C. for 2 minutes, and once at 72° C. for 10 minutes. The PCR product was ligated to a vector pCR2.1 (from Invitrogen) using Takara ligation kit (from Takara Shuzo) and transformed into an Escherichia coli TB1 strain. As a result of determining the entire nucleotide sequence of the PCR product, it consisted of 1029 bp in the length, longer than the reported hrpZ gene (He et al.: Cell 73:1255-1266(1993)) by three bases (one amino acid), and showed a homogoly of 96.7% in nucleotides and a homology of 96.5% in amino acids. The reason that the nucleotide sequences are not completely the same is thought to be due to a variation among the pathover. The nucleotide sequence of the cloned hrpZ gene is shown in SEQ. ID No. 1 and the deduced amino acid sequence obtained therefrom is shown in SEQ. ID No. 2, respectively.

Example 2

Expression in an Escherichia coli and Production of an Antibody

[0042] The above plasmid with an hrpZ gene integrated into pCR2.1 was digested with restriction enzymes BamHI and SaII, and was subjected to electrophoresis on 0.7% agarose to separate a fragment of about 1.1 kb. This fragment was ligated to an expression vector pQE31 (from QIAGEN) digested with the same enzymes and transformed

into Eschrichia coli M15 strain. The thus obtained Eschrichia coli was cultured in an LB medium in the presence of 1 mM of IPTG at 37° C., harpin_{pss} was accumulated as insoluble fraction. Since this protein showed poor adsorption to a nickel resin adsorbent, the purification of harpinpss was conducted in the following procedure. The Eschrichia coli M15 strain having the pQE31 vector with the hrpZ gene integrated thereinto was cultured in 2 ml of an LB medium containing 100 mg/l of ampicillin and 25 mg/l of kanamycin at 37° C. overnight, and transferred into 250 ml of the LB medium and cultured for about three hours; then 1 mM of IPTG was added thereto and the culture was further conducted at 37° C. for 4 hours. Cells were collected by centrifugation, the insoluble fraction was dissolved in 4 ml of an eluation buffer (8 M urea, 0.1 M sodium dihydrogen phosphate, 0.01 M Tris, pH 8.0), and a supernatant liquid was obtained by centrifugation and subjected to electro-phoresis on a 12.5% acrylamide gel containing 0.1% SDS, and then stained with Coomassie Brilliant Blue to cut a band appearing at around 40 kDa. The gel was cut into small pieces, and an elution buffer (1% SDS, 0.02 M Tris HCl, pH of 8.0) was added thereto in an amount ten times the volume of the gel, and shaken for three days. The supernatant was transfered to a dialysis membrane with a cut off molecular weight of 6,000 to 8,000, and the dialysis was conducted with 80% acetone as an external liquid once for 4 hours and once overnight. The whole content in the dialysis tube was moved into an Eppendorf tube, subjected to centrifugation to discard the supernatant, and the pellet was dried to obtain a purified harpin_{pss} preparation. 3 mg of the purified harpin_{pss} was sent to Sawady Technology for the production of an antibody (anti-rabbit harpin_{pss} serum).

Example 3

Construction of a Gene and Transformation of a Plant

[0043] The hrpZ gene integrated into pCR2.1 was excised from the vector by digestion with restriction enzymes Xbal and SacI (from Takara Shuzo). On the other hand, superbinary vector pSB21 (35S-GUS-NOS, Komari et al.: Plant J. 10: 165174 (1996)) was digested with the same enzymes to remove the GUS gene, and the hrpZ gene was integrated thereinto. According to the above procedure, a construct named 35S-hrpZ (35S promoter-hrpZ gene-NOS terminator) was constructed. The cauliflower mosaic virus 35S promoter is a promoter capable of constitutively promoting a high expression, and it is anticipated that rice and tobacco transformed with this construct will accumulate harpin_{pss}, the hrpZ gene product, in the whole body.

[0044] pSB21 was digested with restriction enzymes HindIII and XbaI to remove the 35S promoter, and a 0.9 kb fragment of corn PPDK promoter (Taniguchi et al.: Plant Cell Physiol. 41: 42-48 (2000)) was integrated thereinto. The resulting plasmid was digested with XbaI and SacI to remove the GUS gene, and then the above-described hrpZ XbaI-SacI fragment was inserted thereinto. Thus, PPDK-hrpZ (PPDK promoter-hrpZ gene-NOS terminator) was constructed. The corn PPDK promoter is a promoter capable of promoting a strong expression in photosynthesis organs such as mesophyl cells (Taniguchi et al.: Plant Cell Physiol. 41: 42-48 (2000)), and it is anticipated that rice plants transformed with this construct will accumulate harpin_{pss}, the hrpZ gene product, in green organs (leaves).

[0045] PAL promoter was cloned as below. Plasmid DNA was extracted from agrobacterium LBA4404 strain (gifted from Prof. Shiraishi of Okayama University) having a construct containing PSPAL1 (PSPAL1 promoter-GUS gene-NOS terminator) (Yamada et al.: Plant Cell Physiol. 35: 917-926 (1994), and Kawamata et al.: Plant Cell Physiol. 38: 792-803 (1997)). On the other hand, a reverse primer and two forward primers were designed on the basis of the nucleotide sequence of the reported PSPAL1 promoter (Patent: JP 1993153978-A 1 22-Jun.-1993; TAKASAGO INTERNATL. CORP.):

PALRVXba:
GGG GTC TAG AAT TGA TAC TAA AGT AAC TAA TG
PALFFHIN:
TTG GAA GCT TAG AGA TCA TTA CGA AAT TAA GG
PALFSHIN:
CTA AAA GCT TGG TCA TGC ATG GTT GCT TC

[0046] A promoter region (PAL-S) of about 0.45 kb in the upstream of the starting point of translation (about 0.35 kb at the upstream of the initiation point of transcription) was amplified by the combination of PALRVXba and PALF-SHin, and a promoter region (PAL-L) of about 1.5 kb by the combination of PALRVXba and PALFFHin. The abovementioned agrobacteruium plasmid DNA was used as a template and PCR was conducted with these primers. The reaction conditions of PCR were as below: reaction solution: 50 μ l; each primer: 0.5 μ M, dNTP: 0.2 mM; 1×ExTAq buffer, ExTAq DNA polymerase (from Takara Shuzo): 1U; and the reaction was conducted once at 94° C. for three minutes, then 30 cycles at 94° C. for one minute, at 50° C. for one minute and at 72° C. for two minutes, and once at 72° C. for 6 minutes. A PCR product was cloned to vector pCR11 (from Invitrogen).

[0047] Since the PsPAL1 promoter had a HinIII site at the upstream 142 bp from the starting point of translation, PAL-S was digested completely with restriction enzyme XbaI and then partially with HindIII to obtain a 0.45 kb of fragment from pCR11. The above mentioned pSB21 was digested with HindIII and XbaI to remove the 35S promoter, and PAL-S was integrated thereinto. In the pSB21 vector employed here the unique PvuII site existing in the basic structure had been removed, and, instead, a PvuII linker had been placed at the unique ECOR1 site (just after the Nos terminator). The plasmid with PAL-S integrated thereinto was further digested with XbaI and SacI to remove the GUS gene, and then the above mentioned 1.1 kb hrpZ XbaI-SacI1 fragment was inserted therein. PALS-hrpZ was constructed according to the above procedure. Next, PAL-L integrated into pCR11 was digested with restriction enzymes XhoI and XbaI to take out a 1.45 kb PAL promoter, which was integrated into vector pSB11 (Komari et al.: Plant J. 10: 165-174 (1996)) co-digested with the same enzymes. The formed plasmid was digested with XbaI and SmaI, and an XbaI-PvuII fragment of PALS-hrpZ (hrpZ-NOS terminator) was inserted therein. In this manner, PALL-hrpZ was produced. The PAL promoter promotes a low-level expression constitutively, but it is a promoter strongly induced with a pathogen and an injury (Yamada et al.: Plant Cell Physiol. 35: 917-926 (1994), and Kawamata et al.: Plant Cell Physiol. 38: 792-803 (1997)), and it is anticipated that a tobacco plant transformed with PALS-hrpZ or PALL-hrpZ accumulates more harpin_{pea} at the place of stress when these stresses occur. In this case, it is anticipated that more harpings will accumulate in the case of PALL relative to the case of PALS.

[0048] According to the tri-parental mating system, of Escherichia coli LB392 strain containing the thus produced four constructs 35S-hrpZ, PALS-hrpZ, PALS-hrpZ and PALL-hrpZ (summarized in FIG. 1), agrobacterium LBA4404 strain containing a vector pSB4U with a selection marker gene integrated thereinto (corn ubiquitin promoterhygromycin-resistant gene (hptII)-NOS terminator) and Escherichia coli HB101 containing a helper plasmid pRK2013, the hrpZ gene containing construct was introduced into an agrobacterium utilizing homologous recom-

[0049] The transformation of a tobacco was performed by the leaf disc method (Horsch et al.: Science 227: 1229-1231 (1985)). A leaf of tobacco variety SR1 grown in a greenhouse was sterilized by treatment with ethanol for 30 seconds and with antiformin diluted 5 times for 5 minutes, and after it was cleaned with sterilized water twice, it was cut into one-centimeter squares, and an agrobacterium suspension was inoculated thereto. The concentrations of hygromycin at the time of induction and selection of a transfected shoot and at the time of rooting were 50 or 100 mg/ml and 0 or 50 mg/ml, respectively. For the transformation of rice, immature-embryo-derived cali of varieties of paddy rice, Tsukinohikari, and Koshihikari were transformed employing agrobacterium according to the method of Hiei et al.: Plant J. 6: 271-282 (1994).

Example 4

Analysis of Transformants

[0050] (1) Transgenic Tobacco

[0051] 15 individuals of the re-generated plant were obtained from 35S-hrpZ, 10 individuals were from PALShrpZ and 16 individuals were from PALL-hrpZ. There was observed no remarkable difference between the constructs in transformation efficiency. Western analysis was performed on the primary generation (To) of the transformant, and Western analysis and disease assays were performed on the self-pollinated next generation (T_1) .

[0052] 1) Western Analysis of To Generation

[0053] 2x2 cm of a leaf of a transgenic tobacco of the 4 or 5 leaf stage and 2×2 cm of a leaf of a non-transgenic tobaco (SR1) were pulverized in 0.1 M HEPES-KOH pH 7.6 buffer in a mortar. The supernatant liquid after centrifugation with 15000 g for 10 minutes was made a protein sample. The amount of the protein was determined with a Bio-Rad Protein Assay kit (from BIO-RAD). About 20 µg of the protein was fractioned by the SDS-PAGE method according to the method of Laemmni et al. (Nature 227: 680-685 (1970)), on 12.5% PAGEL (from ATTO). After electrophoresis, the protein bands on the gel were transferred to a PVDF membrane (from Millipore). The PVDF membrane was placed in a 1xTBS buffer containing 0.5% skim milk for 30 minutes, and shaken in the same buffer containing 1/1000 (v/v) of anti-harpin_{pss} serum at room temperature overnight. As a secondary antibody was employed an anti-goat rabbit IgG peroxidase labeled conjugate (from MBL) or an antigoat rabbit IgG alkaline phosphatase conjugate (from BIO- RAD) at the concentration of 1/1000 (v/v). As color development systems were employed HRP Color Development Reagent (from BIO-RAD), alkaline phosphatase substrate kit II (from Vector Laboratories). The amounts of the protein expressed were calculated by comparison with the color development of the harpings sample of a known concentration, by using a densitometer (model GS-670, from BIO-RAD). Some of the results of the Western analysis of the To generation is shown in FIG. 2, and the whole results are summarized in Table 1.

[0054] The expression level is shown in four stages (+++, ++, +, -), which show 0.1% or more of the total soluble proteins (+++), 0.05 to 0.1% (++), 0.05% or less (+) and below the detection limitation (-) in the amount of expression, respectively. This is true also in Tables 2, 3 and 4 to be described later.

TABLE 1

Results of the Western Analysis of the Tobacco To Generation												
	Number of re-generated	Expression level of Harpings										
Construct	individuals	-	+	++	+++ _p							
PALS-hrpZ	10	1	8	1	0							
PALL-hrpZ	16	2	10	4	0							
35S-hrpŽ	15	6	2	1	6							
SR1		3	0	0	0							

*Each numerical value shows the number of individuals showing each

expression level.

The expression level of harpin_{pss} is shown in four stages (+++: particularly high expression, +-: high expression, +-: moderate to poor expression, --: below the detection limitation).

[0055] In the case of the constructs having a PAL promoter, the accumulation of harpin_{pss} was detected in 80% or more of individuals. As anticipated, PALL had a larger proportion of high-expression individuals (++) than PALS. On the other hand, in the case of the construct having a 35S promoter, though no accumulation of harpin_{pes} was detected in 6 individuals of the 15 individuals, high-expression individuals were obtained in 7 individuals, near half of the total individuals. Besides, a very high expression (+++) was shown in 6 individuals. Interestingly, no morphological change was observed in the organ of any of a leaf, a stem, a root or a flower of these high-expression individuals, and seed fertility was normal in almost all of them.

[0056] 2) Western Analysis of the T₁ Generation and Disease Resistance Assay

[0057] Reaction to powdery mildew fungus (Erysiphe cichoracearum) was analized in about 8 lines of KH1-2 (PALS-hrpZ), KC6-7 (PALL-hrpZ), KC8-1 (PALL-hrpZ), KK1-1 (35S-hrpZ), KK3-8 (35S-hrpZ), KK4-2 (35S-hrpZ), KK4-3 (35ShrpZ), KK7-6 (35S-hrpZ), in which the amount of harpin_{pss} accumulated was high in the To generation.

[0058] Tobacco individuals in which harpin pss was accumulated at a high level in the To generation were selected, and seeds of self-pollinated next generation (T1) thereof were obtained. The seeds were sowed and observed for about two months, but no visual morphological change was observed for this period; they grew normally in the same manner as the To generation, and no hypersensitive response was observed on the surface of a leaf. Then, powdery mildew fungi were sprayed to inoculate upon the T1 generation of the transgenic tobacco of the 4 or 5 leaf stage and a disease resistance assay was performed. About 2 L of a suspension of powdery mildew fungi spores (1.4×106 spores/ml) was spray-inoculated to 244 recombinants and 41 original individuals. As a result, hypersensitive-responselike localized necrosis spots were induced onto a lower leaf of the recombinant 4 or 5 days after inoculation (FIG. 3A, B). Surprisingly, not only in the case of the PAL-hrpZ constructs but also in the case of the 35S-hrpZ constructs employing a constitutive promoter, specific localized necrosis spots were induced after the pathogen infection (FIG. 3B). The expression frequency of localized necrosis spots on the 5th day after the inoculation was about 5% in the non-transformants, but the frequency was from 6 to 14 times grater in the 35S-hrpZ construct (30 to 71%), from 4 to 5 times greater in the PAL-hrpZ constructs (20 to 27%) (Table 2), and thereafter, in the case of the PAL-hrpZ constructs, the number of local necrosis spots gradually increased. This was assumed to be due to the response of the PsPAL1 promoter to Erysiphe cichoracearum. Though the amount of harpinpes accumulated and the degree of the formation of localized necrosis spots tended to be positively correlative (Table 3), there were some exceptional transformants in which no accumulation of harpin_{pes} was detected at least in our West-ern analysis but localized necrosis spots occurred.

[0059] Next, in order to examine whether the localized necrosis spots having occurred after the powdery mildew infection were related to disease resistance, the symptom of powdery mildew on the 11th day after the inoculation thereof was examined. As a result, while there existed no individual in which the spread of powdery mildew hyphae was prevented in the non-transformants, from 15 to 57% individuals in the case of 35S-hrpZ constructs and from 13 to 18% individuals in the case of PAL-hrpZ constructs showed apparently less significant symptom as compared to the non-transformants (FIG. 4, Table 2). The prevention of that the spread of powdery mildew was observed not only in leaves with localized necrosis spots but also in middle or upper leaves with no localized necrosis spots, and this is thought to be due to systemic acquired resistance (SAR). As

a result of observing the hyphae of powdery mildew by cotton blue dyeing, the hyphae of powdery mildew extended sharply and spread around the surface in infested leaves of the SR1 of the original line as a control, whereas, though haustorium is formed on the surface of a leaf in the transformants, the spreading of hyphae was prevented and stopped balfway. The promoters employed in the present studies are 35S promoter (constitutive) and PAL promoter (inducible); and it was found that when 35S promoter was employed instead of PAL promoter, the frequency of localized necrosis spots was higher, and it was further found that at least according to examination on the 11th day after inoculation, more individuals with a strong disease resistance were obtained (Table 2). However, it was observed that, in the case of employing the 35S promoter, the localized necrosis spots formed in response to the pathogen became larger (occupying 10% or more of the leaf area) in some individuals, and as a result, lower leaves died out. In addition, inversely, in some individuals with harpinpes accumulated therein, localized necrosis spots were not observable by the naked eye (Table 2), but some of such individuals had resistance to powdery mildew (of individuals with - of localized necrosis spots in Table 2, individuals of the number in parentheses; the amount of harpin expressed is ++ in all). This is thought to be probably due to the occurrence of a hypersensitive response in very small range, and it is possible that a disease-resistant plant with a high practicability can be obtained by the selection of such individuals. According to the fact that no localized necrosis spot occurred without the invasion of the pathogen even in the case where the transription of hrpZ gene was controlled with a constitutive promoter, it is possible to deduce that, since harpin was recognized on the outside of a transmembrane or cell wall of plant cells, probably harpin accumulated in cytoplasm was not recognized for plant cells till the degradation of cells due to the invasion of the fungi, and as a result, it caused a hypersensitive response after the inoculation of the pathogen. Another possibility may be that the elicitor activity of harpin requires the existence of some other factors derived from the pathogen or the plant, induced by the inoculation of the pathogen.

TABLE 2

Relationship among the Amount of harpin_{pss} Accumulated, the Formation of Localized Necrosis Spots and Disease Resistance of the Tobacco T_1 Generation

Line Name	Construct	Expression level (T_0)	Number of individuals analyzed (T ₁)
KH1-2	PALS-hrpZ	++	18
KC6-7	PALL-hrpZ	++	43
KC8-1	PALL-hrpZ	++	44
KK1-1	35S-hrpZ	+++	23
KK3-8	35S-hrpZ	+++	33
KK4-2	35S-hrpZ	++	35
KK4-3	35S-hrpZ	+++	7
KK7-6	35S-hrpZ	+++	41
SR1	(control)	-	41

TABLE 2-continued

Relationship among the Amount of $harpin_{pes}$ Accumulated, the Formation of Localized Necrosis Spots and Disease Resistance of the Tobacco T, Generation

	lo (Nu	ocalized mber of	individu necrosis f individu s of disea	spots	Rate of individuals with localized necrosis spots (5th day after	Rate of individuals with less progress of disease spots (11th day after		
Line Name	+++	++	+	_•	inoculation)	inoculation)		
KH1-2(PALS)	0	0	5(3)	13(0)	27%	16%		
KC6-7(PALL)	0	1(1)	8(6)	34(1)	20%	18%		
KC8-1(PALL)	0	1(0)	11(5)	32(1)	27%	13%		
KK1-1(35S)	0	0	7(3)	16(1)	30%	17%		
KK3-8(35S)	0	2(0)	11(5)	20(0)	39%	15%		
KK4-2(35S)	1(1)	4(3)	15(6)	15(0)	57%	28%		
KK4-3(35S)	0	3(3)	2(1)	2(0)	71%	57%		
KK7-6(35S)	1(1)	4(4)	18(4)	18(1)	56%	24%		
SR1 (control)	0 0 2(0) 39(0)		5%	0%				

The degree of localized necrosis spots is shown in four stages (+++: very high, ++: high,

[0060]

TABLE 3

many transfer and the second control of the second
Relationship between the Expression level of Harpinges and the Number
of Localized Necrosis Spots in the Tobacco T. Generation

Expression level of harpin _{pss} *	Degree of	Incidence of localized necrosis			
(Western analysis)	+++	++	+	_	spots
+++	1	4	19	19	56%
++	0	5	32	77	32%
+	1	6	18	38	40%
-	0	1	5	18	25%
SR1	0	0	2	39	5%

The expression level of harpinpss is shown in four stages (+++: particularly high expression, ++: high expression, +: moderate to poor expression, pelow the detection limit) (SR1, -).

[0061] (2) Transgenic Rice

[0062] 1) Western Analysis of the To Generation

[0063] Harpin was introduced into a rice variety, Tsukinohikari. 35 individuals of the regenerated plant were obtained from the 35S-hrpZ construct, and 26 individuals of the regenerated plant were obtained from the PPDK-hrpZ construct. There was observed no remarkable difference between the constructs in transformation efficiency. Western analysis was performed on the primary generation (To) of the transformation and individuals with a high expression were selected.

[0064] Protein was extracted from the regenerated transgenic rice (Tsukinohikari) in the same manner as in the example of the tobacco and subjected to Western analysis. The results of Western analysis of the To generation are shown in Table 4.

TABLE 4

Results of the Western Analysis of the To Generation of Rice

	Number of regenerated	Expression level of harpings							
Construct	individuals	-	+	++	+++				
35S-hrpZ PPDK-hrpZ	35 26	17 .9	5 13	13 4	. 0				

^{*}Each numerical value shows the number of individuals showing each

[0065] In the case of the rice (Tsukinohikari), similar to the case of the tobacco, individuals with a high-expression of harpin_{pes} were obtained (see also FIG. 2). In the case of a construct having a 35S promoter, the accumulation of harpin_ was detected in about half of the individuals, and the rate of high-expression individuals (++) was about one-third or more of the whole. Also, in the case of a PPDK promoter the accumulation of harpinpss was detected in about twothirds of the individuals, and of them, 4 individuals showed a high expression. Interestingly, no morphological change was observed in the organ of any of a leaf, a root or a flower of these high-expression individuals. And seed fertility was normal in almost all of them, and T1 seeds of high-expression individuals could be obtained.

[0066] 2) Western Analysis of the To Generation and the Disease Resistance Assay of the T₁ Generation

[0067] Next, harpin_{pes} was introduced into Koshihikari, one of the most important varieties of rice of Japan. The results of the Western analysis of the To generation are shown in Table 5.

The degree of localized necrosis spots is shown in four stages (+++: great many, ++: many, +: few, -: nil).

expression level.
The Expression level of harpin_{pes} is shown in four stages (+++: particularly high expression, ++: high expression, +: moderate to poor expression, -: below the detection limit).

TABLE 5

Results of the V	Western Analysis of the T _o Generation of Rice (Koshihikari)
	(Koshihikari)

	Number of regenerated	Expression level of harpings							
Construct	individuals	-	+	++	+++ ^b				
35S-hrpZ PPDK-hrpZ	78 27	18 7	33 13	21 7	6 0				

^{*}Each numerical value shows the number of individuals showing each

[0068] Of the individuals of the T_0 generation with the 35ShrpZ construct introduced thereinto, four individuals house were set at 25° C. under light conditions for 16 hours, and at 22° C. under dark conditions for 8 hours. The evaluation of disease resistance was performed by visually counting the number of progressive disease spots on the 5th leaf at 6th day after the inoculation, said leaf being the topmost development leaf at the time of inoculation. Significant differences among the results were evaluated according to the Mann-Whitney U test.

[0069] As a result, though no localized necrosis spot due to the inoculation of the blast fungi was observed, the average number of progressive disease spots was reduced by 24 to 38% relative to the control Koshihikari in three lines (hrp5-8, hrp42-9, hrp23-5) out of the four lines of the harpin_{pss}-introduced rice. Moreover, this reduction was statistically significant (Table 6). The above results show that the disease resistance of rice could be increased by the introduction of harpin_{pss}.

TABLE 6

Results of the Disease Test against Rice Blast of the Four Lines of Harpin _{pss} - Intorduced Rice (T ₁ Generation)											
Strain	Number of tested individuals	Number of average progressive disease spots ^a (standard error)	pots ^e Significant Test ^b								
hrp5-8	16	9.3 (±1.0)	significant								
hrp23-5	21	11.4 (±1.3)	(significance level 1%) significant (significance level 5%)								
hrp24-1	20	14.4 (±1.4)	No significant difference								
hrp42-9	14	9.4 (±1.4)	significant (significance level 1%)								
Koshihikari	64	15.0 (±0.7)									

[&]quot;Results of the 5th leaf on the 6th day after inoculation

showning a large amount (+++ in Table 5) of the accumulation of harpin_{pss} (hrp5-8, hrp23-5, hrp24-1, hrp429) were selected, and their vulnerability to rice blast in the T, generation was examined. The seed fertility of the selected four high-expression individuals was normal, and many self-fertilized seeds could be obtained. T₁ seeds were sowed in a seedling case with culture soil in a manner of 8 seeds×2 rows, cultivated in a greenhouse, and subjected to a disease assay at the 4.8 to 5.2 leaf stage. As a rice blast fungus (Magneporthe grisea) was employed race 007. For inoculation, a conidium formed by culturing the blast fungi on an oatmeal sucrose agar medium at 28° C. under dark condition and then, after the spread of the fungi, at 25° C., irradiating near ultraviolet light for three days was employed. The inoculation of the blast fungi was performed by sprayinoculating 30 ml of a suspension adjusted to 1.5×10⁵ condia/ml in 0.02% Tween 20 per three seedling cases. The spray-inoculated rice was held in a moistening incubator (SLPH-550-RDS, manufactured by Nippon Medical & Chemical Instruments Co. Ltd.) for 24 hours after the inoculation at 25° C. at a humidity of 100%, and then transferred into the greenhouse. The conditions of the green[0070] As a result of the present invention, it has become apparent for the first time that disease resistance can be imparted to a plant by connecting a gene enconding harpin to a constitutive promoter or an inducible promoter and introducing the gene into the plant. This harpinin-introduced plant is thought to be useful for explicating the function of harpin as a protein elicitor, and also for explicating the mechanism of localized or systemic acquired resistance. In addition, it is revealed that the production of a harpinintroduced resistant plant, which has been thought to be difficult without the use of an inducible promoter, can sufficiently be realized by employing a constitutive promoter, and the extension of the application range of the present approach can be shown. The present invention shows that a method for producing a disease-resistant plant by integrating a DNA sequence encoding a harpin into an expression cassette comprising a sequence of an appropriate constitutive, or organ- or phase-specific promoter capable of functioning in a plant cell, or a promoter induced with stress or pests, and a sequence of a terminator capable of functioning in a plant cell, and introducing it into the plant cell to obtain a regenerated individual is a useful and effective approach in view of genetic engineering.

expression level.

The expression level of harpings is shown in four stages (+++: amount of accumulation of 0.5% or more to the total soluble leaf proteins, ++: amount of accumulation of from 0.1 to 0.5%, +: amount of accumulation of from 0.01 to 0.1%, -: below the detection limit).

Significant difference to Koshihikari in the Mann-Whitney U test

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Leu	Thr 130	Lys	Gln	qaA	Gly	Gly 135	Ala	Ser	Phe	Ser	Glu 140	Asp	qaA	Met	Pro	
145					150					155			Ala		160	
Pro	Lys	Pro	Asp	Ser 165	Gly	Ser	Trp	Val	Asn 170	Glu	Leu	Lys	Glu	Авр 175	Asn	
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Ile	Gly	Gln 195	Gln	Leu	Gly	Asn	Gln 200	Gln	Ser	Gly	Ala	Gly 205	Gly	Leu	Ala	
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<220> FEATURE:
<223> OTHER INFORMATION: Synthetic primer derived from the reported
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<210> SEQ ID NO 7
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-continued

- <211> LENGTH: 29
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- <220> FEATURE:
- 223> OTHER INFORMATION: Synthetic primer derived from the reported PSPAL1 promoter

<400> SEQUENCE: 7

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29

- 1. A transgenic, disease-resistant plant which has been transformed with an expression cassette comprising:
 - a promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression; and
 - a gene, under the control of said promoter, encoding an elicitor protein;
 - wherein said plant is capable of effecting the constitutive, inducible, or organ- or phase-specific expression of the elicitor protein in an amount effective for inducing a defense reaction.
- 2. A transgenic, disease-resistant plant as claimed in claim 1, wherein said promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression and said gene, under the control of said promoter, encoding an elicitor protein, are integrated into the genome.
- 3. A transgenic, disease-resistant plant as claimed in claim 1 or 2, wherein said elicitor protein is a protein possessing a hypersensitive-response-inducing activity against disease microorganisms.
- 4. A transgenic, disease-resistant plant as claimed in claim 3, wherein said protein possessing a hypersensitive-response-inducing activity is selected from:
 - (a) a protein consisting of the amino acid sequence of SEQ. ID No. 2;
 - (b) a protein consisting of an amino acid sequence derived from the amino acid sequence of SEQ. ID No. 2 by deletion, substitution, addition or insertion of one or more amino acids, and possessing a hypersensitiveresponse-inducing activity; and
 - (c) a protein consisting of an amino acid sequence being at least 50% homologous to the amino acid sequence of SEQ. ID No. 2, and possessing a hypersensitive-response-inducing activity.
- 5. A transgenic, disease-resistant plant as claimed in claim 2, wherein said gene encoding an elicitor protein is selected from:
 - (a) a DNA molecule consisting of the nucleotide sequence of SEQ. ID No. 1;
 - (b) a DNA molecule consisting of a nucleotide sequence derived from the nucleotide sequence of SEQ. ID No. 1 by deletion, substitution, addition or insertion of one or more nucleotides, and encoding a protein possessing a hypersensitive-response-inducing activity;
 - (c) a DNA molecule consisting of a nucleotide sequence being hybridizable with a DNA molecule consisting of the complementary nucleotide sequence to the nucleotide sequence of SEQ. ID No. 1 under stringent

- conditions, and encoding a protein possessing a hypersensitive-response-inducing activity; and
- (d) a DNA molecule consisting of a nucleotide sequence being at least 50% homologous to the nucleotide sequence of SEQ. ID No. 1, and encoding a protein possessing a hypersensitive-response-inducing activity.
- 6. A method for producing a transgenic, disease-resistant plant capable of effecting a constitutive, inducible, or organor phase-specific expression of an elicitor protein in an amount effective for inducing a defense reaction, comprising the steps of:
 - (a) obtaining a transgenic plant cell with an expression cassette comprising a promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression and a gene, under the control of said promoter, encoding an elicitor protein; and
 - (b) reconstructing, from said transgenic plant cell, a complete plant.
- 7. An expression cassette for producing a transgenic, disease-resistant plant capable of effecting a constitutive, inducible, or organ- or phase-specific expression of an elicitor protein in an amount effective for inducing a defense reaction, comprising at least:
 - (a) a promoter capable of promoting a constitutive, inducible, or organ- or phase-specific gene expression; and
 - (b) a gene, under the control of said promoter, encoding the elicitor protein.
- 8. An expression cassette as claimed in claim 7, wherein said elicitor protein is a protein possessing a hypersensitive-response-inducing activity against disease microorganisms.
- 9. An expression cassette as claimed in claim 8, wherein said protein possessing a hypersensitive-response-inducing activity is selected from:
 - (a) a protein consisting of the amino acid sequence of SEQ. ID No. 2;
 - (b) a protein consisting of an amino acid sequence derived from the amino acid sequence of SEQ. ID No. 2 by deletion, substitution, addition or insertion of one or more amino acids, and possessing a hypersensitiveresponse-inducing activity; and
 - (c) a protein consisting of an amino acid sequence being at least 50% homologous to the amino acid sequence of SEQ. ID No. 2, and possessing a hypersensitive-response-inducing activity.
- 10. An expression cassette as claimed in claim 7, wherein said gene encoding an elicitor protein is selected from:

- (a) a DNA molecule consisting of the nucleotide sequence of SEQ. ID No. 1;
- (b) a DNA molecule consisting of a nucleotide sequence derived from the nucleotide sequence of SEQ. ID No. 1 by deletion, substitution, addition or insertion of one or more nucleotides, and encoding a protein possessing a hypersensitive-response-inducing activity;
- (c) a DNA molecule consisting of a nucleotide sequence being hybridizable with a DNA molecule consisting of the complementary nucleotide sequence to the nucleotide sequence of SEQ. ID No. 1 under stringent conditions, and encoding a protein possessing a hypersensitive-response-inducing activity; and
- (d) a DNA molecule consisting of a nucleotide sequence being at least 50% homologous to the nucleotide sequence of SEQ. ID No. 1, and encoding a protein possessing a hypersensitive-response-inducing activity.
- 11. An expression cassette as claimed in any one of claims 7-10 for producing a transgenic, systemic acquired disease-resistant plant.
- 12. An expression cassette as claimed in any one of claims 7-11, wherein said elicitor protein is expressed specifically at the time of infection of disease microorganisms in an amount effective for inducing a defense reaction.
- 13. An expression cassette as claimed in claim 12, comprising a constitutive, or organ- or phase-specific promoter.
- 14. A recombinant vector carrying an expression cassette as claimed in any one of claims 7-13.
 - 15. A gene consisting of a DNA molecule selected from:
 - (a) a DNA molecule consisting of the nucleotide sequence of SEQ. ID No. 1;
 - (b) a DNA molecule consisting of a nucleotide sequence derived from the nucleotide sequence of SEQ. ID No. 1 by deletion, substitution, addition or insertion of one or more nucleotides, and encoding a protein possessing a hypersensitive-response-inducing activity;
 - (c) a DNA molecule consisting of a nucleotide sequence being hybridizable with a DNA molecule consisting of the complementary nucleotide sequence to the nucleotide sequence of SEQ. ID No. 1 under stringent conditions, and encoding a protein possessing a hypersensitive-response-inducing activity; and

- (d) a DNA molecule consisting of a nucleotide sequence being at least 50% homologous to the nucleotide sequence of SEQ. ID No. 1, and encoding a protein possessing a hypersensitive-response-inducing activity.
- 16. A gene encoding a protein selected from:
- (a) a protein consisting of the amino acid sequence of SEQ. ID No. 2;
- (b) a protein consisting of an amino acid sequence derived from the amino acid sequence of SEQ. ID No. 2 by deletion, substitution, addition or insertion of one or more amino acids, and possessing a hypersensitiveresponse-inducing activity; and
- (c) a protein consisting of an amino acid sequence being at least 97% homologous to the amino acid sequence of SEQ. ID No. 2, and possessing a hypersensitive-response-inducing activity.
- 17. A protein selected from:
- (a) a protein consisting of the amino acid sequence of SEQ. ID No. 2;
- (b) a protein consisting of an amino acid sequence derived from the amino acid sequence of SEQ. ID No. 2 by deletion, substitution, addition or insertion of one or more amino acids, and possessing a hypersensitiveresponse-inducing activity; and
- (c) a protein consisting of an amino acid sequence being at least 97% homologous to the amino acid sequence of SEQ. ID No. 2, and possessing a hypersensitive-response-inducing activity.
- 18. A transgenic, disease-resistant plant as claimed in any one of claims 1-5, which has been transformed with an expression cassette comprising a constitutive or inducible promoter;

wherein said plant is a transgenic, powdery mildewresistant tobacco.

19. A transgenic, disease-resistant plant as claimed in any one of claims 1-5, which has been transformed with an expression cassette comprising a constitutive promoter;

wherein said plant is a transgenic, blast-resistant rice.

The opine synthase genes carried by Ti plasmids contain all signals necessary for expression in plants

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Signals necessary for in vivo expression of Ti plasmid T-DNA-encoded octopine and nopaline synthase genes were studied in crown gall tumors by constructing mutated genes carrying various lengths of sequences upstream of the 5' initiation site of their mRNAs. Deletions upstream of position -294 did not interfere with expression of the octopine synthase gene while those extending upstream of position - 170 greatly reduced the gene expression. The estimated size of the octopine synthase promoter is therefore 295 bp. The maximal length of 5' upstream sequences involved in the in vivo expression of the nopaline synthase gene is 261 bp. Our results also demonstrated that Ti plasmid-derived sequences contain all signals essential for expression of opine synthase genes in plants. Expression of these genes, therefore, is independent of the direct vicinity of the plant DNA sequences and is not activated by formation of plant DNA and T-DNA border junc-

Key words: Agrobacterium tumefaciens/Ti plasmids/opine synthase genes/promoter regions

Introduction

Crown gall, a neoplastic disease of dicotyledonous plants, develops after infection of wounded tissue with Agrobacterium tumefaciens strains carrying large tumor-inducing (Ti) plasmids (Zaenen et al., 1974; Van Larebeke et al., 1974; Watson et al., 1975). A well-defined segment (T-region) of the Ti plasmid is transferred and covalently integrated, without rearrangements, in plant nuclear DNA (Chilton et al., 1977, 1980; Schell et al., 1979; Thomashow et al., 1980; Lemmers et al., 1980; Zambryski et al., 1980; Yadav et al., 1980; Willmitzer et al., 1980). The transferred DNA (T-DNA) is transcribed (Drummond et al., 1977; Willmitzer et al., 1981a; Gelvin et al., 1981) by the host RNA polymerase II (Willmitzer et al., 1981b).

Transformed crown gall cells are capable of autonomous growth in the absence of exogenous phytohormones (Braun, 1956). Moreover, these plant tumors synthesize a variety of low mol. wt. metabolites (termed opines) which are characteristic for Ti plasmid-induced tumors (Bomhoff et al., 1976), and can be specifically metabolized by agrobacteria growing on the incited tumors (Petit et al., 1970; Petit and Tempé, 1978; Schell et al., 1979; Tempé et al., 1980). The Ti plasmids are currently classified into three groups according to the type of opine they induce in the incited tumors as octopine.

nopaline or agropine Ti plasmids (Guyon et al., 1980).

The T-DNA in octopine tumors consists of two distinguishable segments: TL-DNA and TR-DNA (Thomashow.et al., 1980; De Beuckeleer et al., 1981). TL-DNA, which is essential and sufficient for octopine crown gall formation, codes for eight polyadenylated transcripts, each expressed from an individual promoter (Gelvin et al., 1982; Willimitzer et al., 1982). One of these transcripts (transcript 3) was shown to code directly for the enzyme octopine synthase (Schröder et al., 1981). The nucleotide sequence of this gene was elucidated and both the 5' and the 3' ends of the transcript were precisely identified by S1 nuclease mapping (De Greve et al., 1982). The 5' end of the transcript coding for octopine synthase is located close to the right border of TL-DNA at a distance of 350 – 400 bp. This gene is transcribed from right to left (Willmitzer et al., 1982).

The T-DNA of nopaline Ti plasmids codes for up to 13 polyadenylated transcripts (Bevan and Chilton, 1982; Willmitzer et al., 1983). The region responsible for tumor maintenance is highly homologous between octopine TL-DNA and nopaline T-DNA (Engler et al., 1981). Transcripts and gene functions determined by this conserved 'core' region are common in octopine and nopaline tumors (Joos et al., 1983; Willmitzer et al., 1983). Two different opines were detected in nopaline tumors: agrocinopine (Ellis and Murphy, 1981) and nopaline (Petit et al., 1970). The nopaline synthase gene has been localized by genetic and transcript mapping on HindIII fragment 23 of plasmids pTiC58 and pTiT37 (Holsters et al., 1980; Hernalsteens et al., 1980; Joos et al., 1983; Willmitzer et al., 1983). DNA sequencing of HindIII fragment 23 localized the nopaline synthase gene (Depicker et al., 1982) and the precise position of the right T-DNA borders within HindIII fragment 23 (Zambryski et al., 1982).

To determine whether all signals essential for the expression of the opine synthase genes *in vivo* are located between the 5' initiation site of the opine genes and the junction site with plant DNA or whether expression of these genes is activated by plant DNA sequences, we constructed octopine and nopaline synthase genes with different lengths of sequences upstream of the 5' initiation site and reinserted them in the T-DNA of the Ti plasmids. This approach allowed us to delimit which sequences are important for the *in vivo* expression of the octopine and nopaline synthase genes, and to demonstrate that the plasmid-derived sequences contain all signals necessary for expression in plants.

Results

Expression of the octopine synthase gene in nopaline tumors "Construction of intermediate vectors pGV761, pGV762 and pGV763. The precise number of base pairs in the DNA region between the 5' initiation site of the octopine synthase transcript (De Greve et al., 1982) and the right T-region border sequence (Holsters et al., 1983) has been determined and was found to be 402 (Figure 1a). Therefore, sequences essential for the expression of octopine synthase must either be located in this sequence, or activation of the promoter occurs by junction of the 5' end of the ocs gene with plant

To whom reprint requests should be sent.

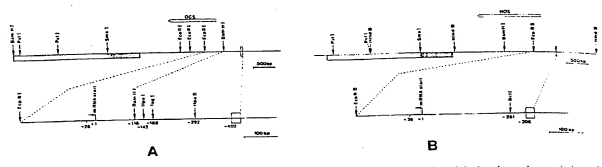


Fig. 1. (A) In the upper part the BamHI fragment 17a and sequences up to the border (white box) are indicated, and the location and transcription polarity of the octopine synthase gene. The white bar shows the homology region between BamHI fragment 17a and the nopaline T-DNA. The hatched portion of the white bar shows the homology region of 750 bp between plasmids pGV761, pGV762, pGV763 and the nopaline Ti plasmid. In the lower part the position of the restriction sites used in this study are indicated with regard to the transcription start of the octopine synthase gene. (B) In the upper part the HindII fragments 23 and 31, and part of the HindIII fragment 22, are indicated (Depicker et al., 1980). The position and transcription polarity of the nopaline synthase gene located in HindIII fragment 23 and the homology region with BamHI fragment 17a of the octopine Ti plasmid pTiAch5 are shown. In the lower part the position of the BcII site is indicated with regard to the transcription start of the nopaline synthase gene.

DNA.

To test which of these possibilities is valid, intermediate vectors containing the octopine synthase gene and different lengths of 5'-flanking sequences (respectively -116 bp, -168 bp and -292 bp from the transcription start; Figure 1a) were constructed and introduced into the nopaline Ti plasmid C58. If the first possibility is correct, these constructions should allow us to delimit the sequences involved in the in vivo expression of the octopine synthase gene. The different steps in the construction of the intermediate vectors are outlined in Figure 2.

Isolation of co-integrated Ti plasmids. As the homology region between plasmids pGV761, pGV762 and pGV763 (Figure 1), and the nopaline Ti plasmid is only 750 bp, we envisaged, to avoid problems of recombination, using the homology of 1270 bp between the amp gene located on pBR322 and the transposon Tn1, inserted into the T-DNA of the nopaline Ti plasmid C58 (Joos et al., 1983; Inzé et al., in preparation).

For this purpose, we selected the plasmids pGV3300 and pGV3305. In pGV3300 a Tn *I* is inserted in *Hind*III fragment 23 just outside the nopaline synthase gene, while in pGV3305 the Tn *I* insertion is located in the nopaline synthase gene. The intermediate vectors pGV761, pGV762 and pGV763 were mobilized from *Escherichia coli* to *Agrobacterium* strains GV3101 (pGV3300) and GV3101 (pGV3305) with the helper plasmids R64*drd*11 and pGJ28 (Van Haute *et al.*, 1983). In all cases, Km^R transconjugants were isolated with a frequency of 10⁻⁸ – 10⁻⁷. Several co-integrate plasmids resulting from a single cross-over were analyzed by DNA/DNA hybridization to confirm their physical structure (data not shown). Recombination always occurred within the homology region common to pBR322 and Tn *I*.

Properties of the co-integrated plasmids. Sunflower hypocotyls and tobacco W38 plants were inoculated with the Agrobacterium strains containing these different co-integrates. The different primary tumor tissues were subsequently analyzed for octopine synthase activity (Otten and Schilperoort, 1978). No octopine synthase activity was detected in sunflower and tobacco tumors induced by the Agrobacterium strains containing the co-integrated plasmids pGV2290 (pGV3300::pGV761) and pGV2291 (pGV3305:: pGV761). Furthermore, in tumors induced by Agrobacterium strains containing the co-integrated plasmids pGV2292

(pGV3300::pGV762) and pGV2293 (pGV3305::pGV762), again no detectable octopine synthase activity could be detected. On the contrary, in sunflower and tobacco tumors induced with *Agrobacterium* strains containing the cointegrated plasmids pGV2294 (pGV3300::pGV763) and pGV2295 (pGV3305::pGV763), octopine synthase activity was detected (Figure 3). The level of activity in these tumors was equal to that found in tumors induced by the *Agrobacterium* strain C58C1 containing an octopine Ti plasmid (pTiB6S3Tra^C).

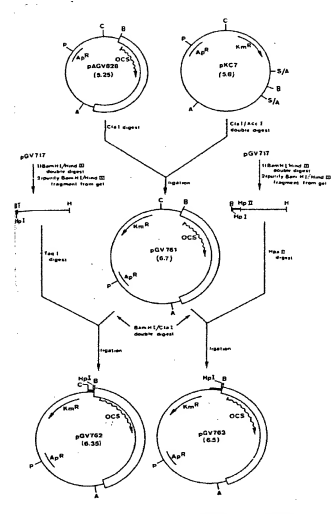
Expression of the nopaline synthase gene in octopine tumors

We have studied the expression of the nopaline synthase gene by a similar approach. DNA sequence analysis showed that the nopaline synthase gene is entirely encoded by the *HindIII* fragment 23 of pTiC58 (Depicker et al., 1982). Furthermore, genomic blotting analysis of nopaline tumor tissue (Lemmers et al., 1980) showed that this *HindIII-23* fragmen is a border fragment. Genomic clones isolated from differen nopaline tumor tissues (Zambryski et al., 1980, 1982; Hol sters et al., 1982) allowed us to determine the exact end point of the T-DNA in crown gall lines. The right T-DNA/plan DNA border is located only 305 bp (Figure 1b) from the star of the nopaline synthase transcript (Depicker et al., 1982).

Construction and properties of pGV2253 and pGV2254

Construction of intermediate vectors pGV705 an pGV706. To demonstrate that the expression of the nopalir synthase gene is independent of the formation of a junctic to plant DNA sequences, and that all sequences involved the in vivo expression of the nopaline synthase gene are pr sent between the start of the mRNA and the end of th T-DNA, we constructed an intermediate vector in which tl sequences between the HindIII site and the BcII site (position -261; Figure 1b) of the HindIII fragment 23 have be deleted and replaced by the SmR gene of R702. This substit tion deletes the 22-bp consensus sequence (position -30 Figure 1b) which is found at the ends of nopaline and C topine T-regions, and which might play a key role in the tegration of the T-region into the plant genome (Zambryski al., 1980, 1982; Simpson et al., 1982; Yadav et al., 191 Holsters et al., 1982, 1983). The construction of the int mediate vector pGV705 is shown in Figure 4.

pGV705 consists of EcoRI fragment 12 of pTiAch5 which the internal HindIII-36a fragment has been substitu



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ig. 2. Construction of intermediate vectors pGV762 and pGV763. The ccl-Clal fragment of pKC7 containing the Km gene was ligated to Clalgested pAGV828. After ligation and selection on ApKm plates, recomnants were screened for the orientation of the Km-resistant fragment by whole digestion with Clal and BamHI. A recombinant plasmid pGV761 as digested with BamHI and Clal, and ligated to the purified HindIII-mHI fragment of pGV717, which contains sequences 5' upstream of the mHI site at -116 in the promoter region of the octopine synthase gene igure 1; Holsters et al., 1983), digested with either Taql or HpaII. By eening recombinant plasmids for the presence of a HpaI site (Figure 1), 3V762 and pGV763 were obtained. Abbreviations: A, Accl; B, BamHI; ClaI; H, HindIII; HpI, HpaI; HpII, HpaII; P, PstI; S, SaII; T, TaqI.

the *Hind*III-*Bcl*I fragment of the nopaline *Hind*III fragent 23 joined to the *BamHI-Hind*III fragment of plasmid 1702 containing the Sm^R gene. This *Hind*III fragment inted in the other orientation in the *Eco*RI fragment 12, is lled pGV706.

Isolation of pGV2253 and pGV2254. The intermediate vects pGV705 and pGV706 were mobilized from E. coli to probacterium strain GV3000 carrying a transfer-constitutive iB6S3 plasmid with the help of the plasmids R64drd11 and iJ28 (Van Haute et al., 1983). Streptomycin-resistant Agroterium strains were obtained in both cases with a joint nester and recombination frequency of 10⁻⁶. The Smistant transconjugants were tested directly for Km sensitivi-

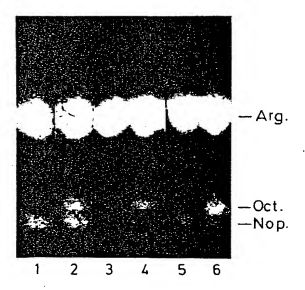


Fig. 3. Detection of octopine in tumors induced with Agrobacterium strains containing the mutant plasmids. 2μ of extracts of tumor tissue before (lanes 1, 3, 5) and after (lanes 2, 4, 6) I h incubation were spotted onto Whatman 3MM paper and subjected to electrophoresis. Lanes 1 and 2: extracts obtained from tissue infected with Agrobacterium containing pGV2295; lanes 3 and 4: extracts obtained from tissue infected with Agrobacterium containing pGV2294; lanes 5 and 6: extracts obtained from tissue infected with Agrobacterium containing pGV2294.

ty. Three percent of the Sm^R transconjugants were Kmsensitive and were double recombinants. The structure of two plasmids pGV2253 and pGV2254 was confirmed by DNA-DNA hybridization (data not shown).

Properties of pGV2253 and pGV2254. Agrobacterium strain containing either pGV2253 or pGV2254 were used to incite tumors on tobacco plants. These tumors synthesize nopaline and octopine (Figure 3), but no mannopine or agropine could be detected. This observation indicates that the deletion substitution of the small HindIII fragment 36a abolishes the synthesis of mannopine and agropine.

Morover, since the sequences between the end of the nopaline T-DNA (position -305) and the BcII site (position -261) have been deleted and replaced by the Sm^R gene of pR702, the 5'-flanking region of the nopaline synthase gene in this construction is separated from TR sequences located to the right (in pGV2253) or to the left (in pGV2254), by the Sm^R insert fragment. Therefore, all the sequences involved in the *in vivo* expression of the nopaline gene must lie within the 5'-flanking region between the start of transcription and the BcII site (position -261).

Discussion

Most of the understanding of the regulatory events controlling gene expression in higher eukaryotes is derived from studies with animal viruses. Several eukaryotic promoters have been examined both by DNA sequencing and by in vitro and in vivo analysis of mutants. These studies have led to the identification of the so-called Goldberg-Hogness or TATA box, a signal that is involved in the precise positioning of 5' RNA ends of genes transcribed by RNA polymerase II (Breathnach and Chambon, 1981; Shenk, 1981). Although the TATA box seems to be both necessary and sufficient for

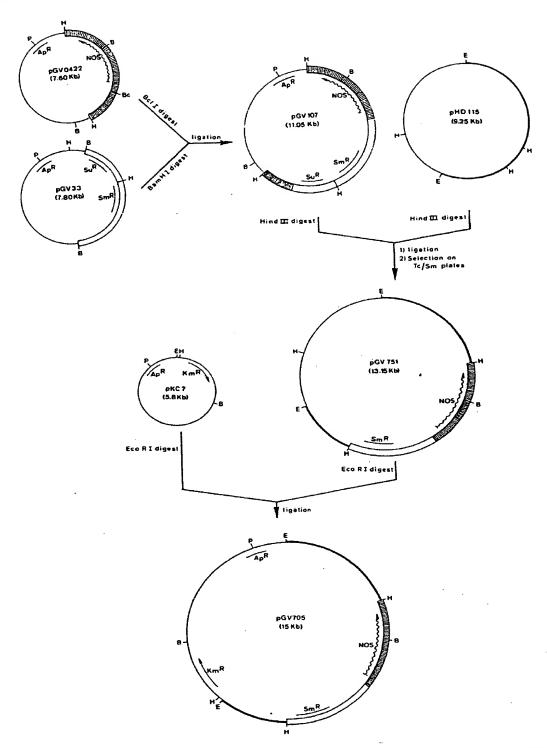


Fig. 4. Construction of the intermediate vector pGV705. Plasmids pGV0422 was linearized with BcII and ligated to BamHI-digested pGV33. After transformation, recombinants were selected on Ap/Sm plates. One of the recombinants, pGV107, was digested with HindIII and ligated to HindIII-digested pHDII5, containing the EcoRI fragment 12 of pTiAch5. After selection on Tc/Sm plates a recombinant, pGV751, was digested with EcoRI and ligated to EcoRI-digested pKC7, making it possible to use the mobilizing method described by Van Haute et al. (1983). Indeed, pGV751, a pACYC184 derivative, cannot be mobilized by pGJ28 and R64drd11.

Table I. Bacterial strains and plasmids

•	Antibiotic resistance	Characteristics	Dimension (kb)	Origin
Strains				
E. coli				
K514		thr leu thi hsdR		Colson et al. (1965)
A. tumefaciens				
GV3101		Rif ^R derivative of C58, cured for pTiC58		Van Larebeke et al. (1974)
GV3105		Ery ^R CmI ^R derivative of C58, cured for pTiC58		Holsters et al. (1980)
Plasmids				
pKC7	Ap Km	HindIII-BamHI of Tn5 in pBR322	5.8	Rao and Rodgers (1979)
pGV0153	Ap	BamHI-8 of pTiAch5 in pBR322	11.6	De Vos et al. (1981)
pGV0201	Ap	HindIII-1 of pTiAch5 in pBR322	16.9	De Vos et al. (1981)
pGV0422	Ap	HindIII-23 of pTiC58 in pBR322	7.6	Depicker et al. (1980)
pGV705	Ap Km Sm	HindIII fragment containing the nos gene and Sm/Sp marker of R702 in EcoRI-12	15	This work
pGV706	Ap Km Sm	HindIII fragment containing the nos gene and Sm/Sp marker of R702 in EcoRI-12, but in opposite direction	15	This work
pGV717	Ap	Hindlil-BamHI fragment of gcl rGV1-1 in pBR322	5.1	Holsters et al. (1983)
pAGV828	Ар	BamHI-Smal of pGV99 in pBR322	5.25	Herrera-Estrella et al. (1983)
pGV761	Ap Km	Clal-Accl of pKC7 in pAGV828	6.7	This work
pGV762	Ap Km	TagI-BamHI of pGV717 in pGV761	6.35	This work
pGV763	Ap Km	Hpall-BamH1 of pGV717 in pGV761	6.5	This work
pGV33	Ap Sm/Sp Su	3.5 kb BamHI fragment of R702 in pBR322	7.7	J. Leemans
pHD115	Tc	EcoRI-12 fragment of pTiAch5 in pACY184	9.25	J. Velten
R702	Km Sm/Sp Tc Su Hg	P-type plasmid	69.0	Hedges and Jacobs (1974)
R64drd11 .	Tc Sm	Ic-type plasmid, transfer-derepressed derivative of R64	109.0	Meynell and Datta (1967)
pGJ28	Km/Nm	Cda ⁺ Ida ⁺ ColD replicon carrying ColE1 mob and bom	9.7	Van Haute et al. (1983)
pGV3100	_	pTiC58, derepressed for autotransfer	212	Holsters et al. (1980)
pGV3300	Ар	pGV3100::Tn <i>1</i>	215	Joos et al. (1983)
pGV3305	Ар	pGV3100::Tn1	215	D. Inzé
pTiB6S3Trac		pTiB6S3, derepressed for autotransfer	192	Petit et al. (1978)

accurate initiation of transcription in vitro (Corden et al., 1980; Wasylyk et al., 1980), regions further upstream are required for efficient in vivo transcription (Grosschedl and Birnstiel, 1980; Benoist and Chambon, 1980; McKnight et al., 1981; Grosveld et al., 1982; Weiher et al., 1983). Recently, a detailed analysis of the promoter of the herpes simplex thymidine kinase (TK) gene (McKnight and Kingsbury, 1982) resulted in an identification of three essential regions within 105 bp upstream of the RNA initiation site.

In higher plants, on the contrary, little is known about sequence signals controlling gene expression. In octopine and nopaline crown gall tumor tissues, the T-DNA is transcribed by RNA polymerase II (Willmitzer et al., 1981a), and encodes a set of well-defined polyadenylated transcripts. Therefore, the T-DNA genes can serve as models for defining transcriptional and translational control sequences in nuclear, proteincoding plant genes. In a first approach, we have attempted to determine which are the minimal 5' upstream sequences in-

volved in the in vivo expression of these opine genes. Deletion of sequences upstream of position -170 of the octopine synthase gene greatly reduces or abolishes the gene expression, while deletion of sequences upstream of position -294 does not interfere with a wild-type level of gene expression. In this sequence of 125 bp an essential region controlling the expression of the octopine synthase gene might be located. Also in the case of the nopaline synthase gene, the 5' sequences downstream of position -261 contain all the information necessary for the in vivo expression of this gene. Therefore, the estimated maximum size of the octopine and nopaline synthase gene promoters are 295 bp and 261 bp, respectively. Although the DNA sequences directly involved in the expression of the opine synthase genes in plant cells are not defined in this study, and identification of these sequences could help in the elucidation of the mechanisms of plant cellular gene control, the results described above clearly demonstrate that the expression of octopine and nopaline synthase genes is determined directly by their 5' upstream flanking sequences and is independent of the direct vicinity of the plant DNA sequences.

Materials and methods

Bacterial strains and plasmids

Bacterial strains and plasmids are listed in Table 1.

Media and culture conditions

Luria broth (LB) and minimal A (minA) media were as described (Miller, 1972). Nitrogen-free medium for the use of octopine or nopaline as sole nitrogen source were as described (Bomhoff et al., 1976). E. coli cultures were grown at 37°C and A. tumefaciens at 28°C. Antibiotic concentrations used for E. coli and A. tumefaciens were respectively, carbenicillin (Cb), 100 μg/ml; streptomycin (Sm), 20 μg/ml and 300 μg/ml; spectinomycin (Sp), 50 μg/ml and 100 μg/ml; kanamycin (Km), 25 μg/ml; rifampicin (Rif), 100 μg/ml; erythromycin (Ery). 50 μg/ml for Agrobacterium; chloramphenicol (Cml), 25 µg/ml for Agrobacterium.

Plasmid isolation

Plasmids were prepared from E. coli by density gradient centrifugation in a CsCl-ethidium bromide gradient of cleared SDS lysates (Betlach et al., 1976). For screening of recombinant plasmids, plasmid DNA was obtained from 10 ml cultures as described (Klein et al., 1980).

DNA analysis

Restriction enzyme analysis, agarose gel electrophoresis, conditions for DNA ligation and transformation of competent E. coli were as described (Depicker et al., 1980). DNA fragments were extracted from low-gelling agarose gels as described (Wieslander, 1979). Total DNA of Ti plasmidcontaining Agrobacterium strains was prepared, digested, separated on agarose gel, transferred to nitrocellulose paper, and hybridized against radioactively labeled recombinant plasmids as described (Dhaese et al., 1979).

Induction and culture of crown gall tumors

Sterile 1-month-old tobacco plants (Wisconsin 38 or SR1) were decapitated and infected with freshly grown agrobacteria. Three weeks later, tumors were excised and transferred to hormone-free Murashige and Skoog medium (Murashige and Skoog, 1962) containing sucrose (30 g/l) and 0.5 mg/ml HR756 (Hoechst A.G.). The tumor tissues, transferred every month, were usually free of bacteria after three transfers, and were further cultivated on antibiotic-free Murashige and Skoog medium. Sunflower hypocotyl segments were inoculated as described by Petit and Tempe (1978).

Detection of opines in plant tumor tissue

Octopine and nopaline detection. The presence of octopine or nopaline in tumor tissue was tested as described by Leemans et al. (1981). Octopine or nopaline synthase activity were determined in vitro according to Otten and Schilperoort (1978).

Agropine and mannopine detection. Agropine and mannopine were detected in tumor tissue as described by Leemans et al. (1981).

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A plant signal sequence enhances the secretion of bacterial ChiA in transgenic tobacco

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Abstract

When the secreted bacterial protein ChiA is expressed in transgenic tobacco, a fraction of the protein is glycosylated and secreted from the plant cells; however most of the protein remains inside the cells. We tested whether the efficiency of secretion could be improved by replacing the bacterial signal sequence with a plant signal sequence. We found the signal sequence and the first two amino acids of the PR1b protein attached to the ChiA mature protein directs complete glycosylation and secretion of the ChiA from plant cells. Glycosylation of this protein is not required for its efficient secretion from plant cells.

Introduction

In eukaryotes most secreted proteins have been shown to possess a signal sequence of approximately thirty amino acids at the N-terminus, which when recognized by the appropriate cellular machinery leads to the translocation of the protein across the membrane of the endoplasmic reticulum [15]. Signal sequences show little homology at the amino acid level but do share common features including positive charge at the amino-terminus, an internal stretch of hydrophobic amino acids, and a polar carboxy-terminal region which contains the cleavage site [22]. These features are conserved in the eukaryotic and prokaryotic kingdoms with some signal sequences across kingdom boundaries [20, 24].

We have shown previously that when the Serratia marcescens chiA gene (which codes for a secreted protein, ChiA) is expressed in tobacco cells, a fraction of the expressed protein is modified by the attachment of complex glycans and secreted from plant cells [12]. In this paper, we describe experiments directed towards improving secretion of ChiA by plant cells. We have tested whether secretion depends upon the presence of a N-terminal signal sequence and if replacement of the signal sequence of ChiA with that of the tobacco PR1b protein increases secretion of the ChiA protein by plant cells. The secretion of mutated forms of ChiA lacking the consensus sequence for N-linked glycosylation was also investigated.

Materials and methods

Plasmid construction

The pChiA plant transformation series derivatives were all prepared in the binary plasmid pJJ2964. This plasmid contains T-DNA carrying an nptII gene driven by the nos promoter (to enable selection of transformed tissue on kanamycin), and unique Bam HI and Hind III cloning sites. Manipulations on the chi4 gene were carried out with it cloned in the vector pUC118 as a fragment containing the cauliflower mosaic virus (CaMV) 35S promoter followed by a leader from the petunia Cab22L gene [6], upstream from the complete chiA gene from Serratia marcescens. Downstream from the chiA gene is a fragment carrying the polyadenylation signals from the Agrobacterium tumefaciens nopaline synthase (nos) gene. The chiA gene had the following modifications to its sequence [7]: (1) a novel Nco I site at position + 1; (2) a novel Sma I site at position 78; (3) the Sma I site present in the native sequence at position 951 has been removed. Oligonucleotide-directed mutagenesis was used to make all these changes. The novel Nco I site changes the second amino acid in the signal peptide from Arg to Ala; the other changes have no effect on the protein sequence.

To construct the plasmid pChiA, the Bgl II-Hind III fragment from the pUC118 derivative described above was ligated into Bam HI-Hind III-cut pJJ2964. The plasmid pChiA-M was constructed following oligonucleotide loop-out mutagenesis of the chiA gene cloned in pUC118, which removed all the codons of the ChiA signal sequence (amino acids 2 to 23) except for the initiator methionine. The Bgl II-Hind III fragment carrying the modified chiA gene and the plant expression signals was then ligated into pJJ2964 as described for pChiA to form pChiA-M.

The plasmid pPRSSChiA was constructed by synthesizing the codons for the PR1b signal sequence plus the first two amino acids of the mature PR1b protein as two complementary oligonucleotides, with a half Nco I site at the 5' end and a half Sma I site at the 3' end. This was

ligated into the Nco I and Sma I sites at the 5' end of the chiA gene. pPRSSChiA was then constructed by ligating the Bgl II-Hind III fragment into pJJ2964, as described above for the other pChiA plasmids.

To remove the glycosylation sites from the ChiA protein, the codons for amino acids at potential N-glycosylation sites (Asn-X-Ser/Thr) were identified on the DNA sequence, then oligonucleotide-directed mutagenesis was used to change the codon for the third amino acid in each site to alanine. All manipulations were carried out on the chiA gene cloned in pUC118 and all changes were verified by DNA sequence analysis. The plasmids pChiA-G and pPRSSChiA-G were then constructed; these are identical to pChiA and pPRSSChiA except that both contain all four of the site-directed mutations that remove the four consensus glycosylation sites.

Plant cell tissue culture

Plant transformations, establishment, maintenance and sampling of suspension cultures, and protoplast preparations were as described [12]. All plant transformations were carried out using *Nicotiana tabacum* cv. SR1.

Protein extraction and measurement

Protein extractions, electrophoresis and immunoblotting of protein extracts were all carried out as described [12], except that immunoblots were developed using an alkaline phosphatase conjugate in place of the horseradish peroxidase conjugate. The buffer used for making protein extracts for ge and enzyme analysis contained 84 mM sodiun citrate. 32 mM sodium phosphate, 6 mM ascor bic acid, and 14 mM β -mercaptoethanol, pH 5.5

Nucleic acid analysis

DNA manipulations were carried out as described [13] or according to enzyme supplier:

instructions. RNA extraction from leaf tissue and primer extension analysis for the quantification of steady-state RNA and confirmation of transcription start sites was carried out as described [6]. Oligonucleotide-directed mutagenesis was by the method of Kunkel [11]; all changes were confirmed by DNA sequence analysis as described by Sanger et al. [17]. Oligonucleotide primers for mutagenesis and sequence analysis were made on an Applied Biosystems 381A DNA synthesizer.

Results

To test whether the bacterial signal sequence of ChiA is required for plant cell secretion, we prepared a deletion mutant of the chiA gene lacking the region which specifies the codons of the signal sequence, pChiA-M (amino acids 2 to 23); the amino terminal of the resulting protein from pChiA-M is shown in Fig. 1. The ChiA protein was then expressed in plant cells with and without its signal sequence by transformation with the binary plasmids pChiA and pChiA-M. In parallel, to determine whether the fraction of ChiA secreted by tobacco cells could be increased by fusion to the signal sequence from a secreted plant protein, we constructed a translational fusion between PR1b and the mature ChiA protein. We chose the tobacco PR1b protein as the source of a plant signal sequence because complete sequence information was available for the PR1b gene and the extracellular location of the protein has been well studied. The portion of the chiA gene encoding the signal sequence of ChiA was replaced with that encoding the signal sequence from PR1b so that the resulting fusion protein contains the PR1b signal sequence plus the first two amino acids of the PR1b mature protein (Gln-Asn) in place of the first two amino acids of the mature ChiA protein (Ala-Ala) (see Fig. 1). This fusion protein was also expressed in plant cells by transformation with the binary plasmid pPRSSChiA. At least 10 independent transformants were prepared for each of the constructions pChiA, pChiA-M and pPRSSChiA, then 2 plants from each group were selected for subsequent comparative analyses. The representative plants were chosen so that the transformants carrying the different chi.4 genes each were expressing similar steady-state chia mRNA levels and ChiA protein.

Immunoblots of leaf proteins isolated from two plants transformed with pChiA show multiple bands (Fig. 2, lanes 2 and 3), the most prominent of which comigrates with ChiA protein expressed in Escherichia coli (Fig. 2, lane 1). We have previously shown that the most prominent species is an intracellular form, and the fainter bands of higher molecular weight are glycosylated, secreted forms of ChiA [12]. Immunoblots of protein from two plants transformed with pChiA-M (Fig. 2, lanes 6 and 7) show a single band which comigrates with ChiA from E. coli; no species of higher molecular weight can be detected, which indicates that glycosylation of ChiA does not occur when it is expressed without a signal sequence. Immunoblot analysis of leaf extracts from plants transformed with pPRSSChiA shows that, in contrast to those transformed with pChiA, all of the cross-

pChiA

MAKENKPLLA LLIGSTLCSA AQA AAPGKPI

pChiA-M

MAAPGKPT

pprsschia MGFLLFSQMP SFFLVSTLLL FLIISHSSHA ONPGKPT

Fig. 1. Deduced amino acid sequences of the deletion and fusion derivatives of the chi.1 gene. The sequences in each case extend to the 7th amino acid in the mature ChiA sequence. The vertical arrow indicates the likely site of cleavage of the signal peptide (known for ChiA in E. coli and predicted for PRSS on the basis of the known cleavage point in the PR1b protein).

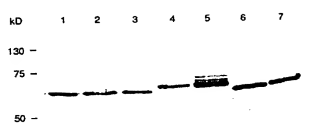


Fig. 2. Immunoblot with Chi.A antibody to the total leaf protein (100 μg) from individual tobacco plants transformed with ChiA derivatives. Lane 1: Chi.A from E. coli (200 ng); lanes 2 and 3: pChiA (ChiA signal sequence): lanes 4 of 5: pPRSS-ChiA (PR1 signal sequence): lanes 6 and 7; pChiA-M (no signal sequence).

reacting protein is in a position corresponding to the glycosylated forms of ChiA (Fig. 2, lanes 4 and 5).

We assayed the level of secretion of ChiA from these different transgenic plant cells by analyzing the media from plant cell suspension cultures and by comparing the profiles of protein extracts from leaf protoplasts and corresponding whole leaves. We have shown that these approaches give consistent results and correctly demonstrate secretion of the PR1b secreted tobacco protein (unpublished data). The culture fluid from suspension cultures established from individual plants transformed with pChiA, pChiA-M or pPRSSChiaA, was analyzed by immunoblotting (Fig. 3). There is little or no ChiA protein in the medium from the pChiA-M transformed cells (lanes 4 and 5), and high levels of ChiA in the medium from the pPRSSChiA or pChiA transformed cells (lanes 2, 3, 6 and 7). Furthermore the ChiA which is present in the culture medium from pPRSSChiA and pChiA transformants is the higher-molecularweight glycosylated form. The faint band seen in lanes 4 and 5, which comigrates with the bacterial standard (lane 1), probably corresponds to non-glycosylated non-secreted ChiA which is in the culture fluid as a consequence of cell death. These data from the analysis of suspension culture media suggest that in the absence of any signal sequence (pChiA-M) the ChiA which is



Fig. 3. Immunoblot with ChiA antibody to protein isolated from suspension culture medium. Lane 1: ChiA protein from E. coli (200 ng); lanes 2 and 3: medium from pPRSSChiA cells; lanes 4 and 5: medium from pChiA-M cells; lanes 6 and 7: medium from pChiA cells (lanes 2–7 each contain protein extracted from 1 ml culture medium); lane 8: 100 μg leaf protein from ChiA plant.

expressed is not secreted. In the presence of signal sequence, either the ChiA signal (pChiA or the PR1b signal (pPRSSChiA), ChiA protei is glycosylated and secreted. The observation the higher levels of ChiA protein appear in the m dium from pPRSSChiA transformants (Fig. lanes 2, 3) than from pChiA transforman (Fig. 3, lanes 6, 7) suggests that secretion is mo efficient when the PR1b signal is fused to Chi

Since secreted proteins will be present in le tissues extracts but absent from washed prot plasts, we have compared these tissues from t transgenic plants to further determine wheth secretion is occurring. The results from typic experiments comparing these two tissues : shown in Fig. 4. In the total leaf extract from plant transformed with pCHiA (lane 7), differ molecular weight forms of the ChiA protein (be seen which correspond to glycosylated (upp and non-glycosylated (lower) forms of the p tein. In washed protoplasts of these plants, c the non-glycosylated (lower) form of ChiA is s (lane 6), which is consistent with our above re indicating that the glycosylated higher-molecu weight forms are secreted from plant cells plants transformed with pChiA-M, the prc profiles of washed protoplasts (lane 2) and 1 leaf extract (lane 3) are identical and corresp to the E. coli standard (lane 1), indicating

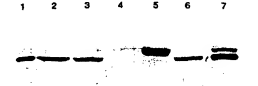


Fig. 4. Immunoblot of protein from leaf or from washed protoplasts. Lane 1: ChiA from E. coli (200 ng); lanes 2 and 3: pChiA-M; lanes 4 and 5: pPRSSChiA; lanes 6 and 7: pChiA. Lanes 2, 4 and 6 are from protoplasts; lanes 3, 5 and 7 are from leaf. Lanes 2 to 7 each contain 100 μg total protein.

glycosylation has not taken place and that little or no secretion is occurring. The profiles from washed protoplasts and total leaf extracts of pPRSSChiA transformants are shown in lanes 4 and 5, and in this case the ChiA protein is present solely as a higher-molecular-weight form, none of which is detected inside washed protoplasts. These data comparing proteins from protoplasts and total leaf extracts confirm that secreted and glycosylated forms of ChiA protein occur only if a signal sequence is attached, and if the PR1b signal sequence is used then all of the ChiA protein is secreted and glycosylated.

There are four consensus N-glycosylation sites (Asn-X-Ser/Thr) in the predicted ChiA protein sequence. We constructed a derivative of the chiA gene where all four sites were 'inactivated' by altering the last codon in the consensus site to Ala. To express the mutant ChiA proteins in plant cells, the binary plasmids pChiA-G and pPRSSChiA-G were used to produce transformed tobacco plants. RNA and protein analysis was used to identify plants expressing high levels of the mutant ChiA proteins. (We noted that expression at both the RNA and protein level was generally significantly higher for plants transformed with pPRSSChiA-G than pChiA-G.)

We compared the protein profile in washed protoplasts with that in total leaf extracts from pChiA-G and pPRSSChiA-G transformants (Fig. 5). The ChiA protein in all plant extracts co-migrated with the mature ChiA protein as purified from bacteria, as would be predicted if no



Fig. 5. Immunoblot of protein from protoplast and leaf extracts from plants expressing ChiA lacking glycosylation sites. Lane 1: E. coli ChiA (150 ng); lanes 2 and 3: pChiA-G; lanes 4 and 5: pPRSSChiA-G. Lanes 2 to 5 each contain 50 μg total protein; lanes 2 and 4 are leaf extracts, lanes 3 and 5 are protoplast extracts.

glycosylation were occurring. In pChiA-G transformants, the intensity of the ChiA band was greater in total leaf extract (lane 2) than in protoplast extracts (lane 3), suggesting that, as with the wild-type protein, secretion of ChiA does occur but not all of the ChiA is being secreted from the cells. In pPRSSChiA-G transformants, a trace of ChiA was detected in the protoplast extracts (lane 5) in contrast to the very high levels in total extract from leaf (lane 4), indicating that most or all of the unglycosylated ChiA protein is being secreted.

Discussion

We have investigated the secretion of the bacterial ChiA protein from plant cells. We had previously demonstrated that ChiA fused to the bacterial signal sequence is inefficiently secreted by plant cells. Here we show that the ChiA protein is fully secreted when the signal sequence derived from the tobacco PR1b protein is fused to the ChiA mature protein and secretion does not occur in the absence of a signal sequence. The lack of secretion in the absence of a signal sequence is expected, given the role of signal sequences in mediating targeting of proteins to the lumen of the endoplasmic reticulum in eukaryotic cells [16].

The fact that no detectable glycosylation of ChiA occurs in the absence of a signal sequence is also expected, since the initial transfer of glycans to proteins occurs as the proteins cross the ER membrane [9].

The most likely explanation for the improved efficiency of ChiA secretion in pPRSSChiA transformants is that the possession of a plant signal sequence improves the ability of the ChiA protein to enter the secretory pathway of the plant cells in which it is expressed. An alternative explanation is that the mature ChiA peptides arising from pChiA- and pPRSSChiA-transformed plants differ in the two N-terminal amino acids, and this difference could alter the mature protein so that it would behave differently in the plant secretory pathway. While this explanation cannot be ruled out, we believe it to be less likely, as we have not detected any significant differences in the physical or enzymological properties of the ChiA expressed with a bacterial or a plant signal sequence. (The precise point of cleavage of the signal sequences when expressed in plant cells remains to be determined.)

Signal sequences show considerable degeneracy, so that even random peptide sequences can function as signal sequences [8]; however, significant differences between prokaryotic and eukaryotic signal sequences are revealed when large numbers of sequences are analyzed statistically [23], These differences may be reflected in the ability of signal sequences to function efficiently in heterologous hosts. There are reports where the use of a signal sequence native to the organism in which the protein is being expressed can enhance the secretion of a heterologous protein [1, 2, 19], and others where more efficient secretion of a foreign protein is seen when it possesses its own signal sequence rather than one derived from the organism in which it is expressed [2, 19]. Determining which features of the PR1b N-terminus are relevant in mediating the efficient secretion of ChiA from plant cells would be an interesting area for further study.

The fact that ChiA is apparently completely located outside the cell when expressed with a plant signal sequence may be taken as further

evidence that the pathway for secretion in plant cells is a default pathway, requiring no positive sorting information other than the possession of a functional signal sequence. Thus it seems likely that many other proteins could also be engineered to be plant secretory proteins. In support of this, Denecke et al. [3] have recently shown that three normally cytoplasmic proteins can be secreted from plant cells by the attachment of a suitable signal sequence.

Glycan side-chains attached to proteins probably have multiple roles [14]; it has often been observed that prevention of glycosylation also prevents the secretion of the altered protein. Sometimes this can be attributed to decreased stability of the altered protein to proteases [4, 14], or to aggregation of the protein [5]. There are also cases where the non-glycosylated forms of the protein are secreted as efficiently as are the glycosylated forms [10, 18]. Thus the role of glycans in intracellular targeting is not simple and cannot be generalized. The likelihood of a direct role for glycan residues in some aspect of protein targeting in eukaryotic cells (for example, by interacting with a receptor as opposed to simply changing the physical properties of the protein) seems remote. Only in the case of lysosome is targeting mediated by mannose-6-phosphate residues [21]. The results presented in this paper demonstrate clearly that the efficiency with which ChiA can be secreted by plant cells is not influenced by the extent to which it is glycosylated.

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